

APPENDICES

**Inventorship Review for U.S. Serial No. 09/930,915
for IMMUNOGENIC HBc CHIMER PARTICLES
HAVING ENHANCED STABILITY; and
U.S. Serial No. 09/931,325 for MALARIA
IMMUNOGEN AND VACCINE**

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:	Ashley J. Birkett)	<u>PATENT</u>
Serial No.:	09/930,915)	Attorney Docket
)	ICC-102.2
Filed:	August 15, 2001)	(4564/81175)
)	
For:	IMMUNOGENIC HBc CHIMER)	
	PARTICLES HAVING ENHANCED)	
	STABILITY)	Group Art No.
)	1648
Examiner:	Donna C. Wortman)	

REPLY AND AMENDMENT

MAIL STOP NON-FEE AMENDMENT
Commissioner for Patents
P.O. Box 1450
Alexandria, Virginia 22313-1450

Sir:

In response to the Action mailed on January 9,
2004, please amend the application as follows.

1. (currently amended) A recombinant chimer hepatitis B core (HBc) protein molecule up to about 515 amino acid residues in length that

(a) contains an HBc sequence of at least about 130 of the N-terminal 150 amino acid residues of the HBc molecule that include a peptide-bonded heterologous epitope or a heterologous linker residue for a conjugated epitope present in the HBc immunodominant loop, ~~or a sequence of at least about 135 residues of the N-terminal 150 HBc amino acid residues,~~

(b) contains one to ten cysteine residues toward the C-terminus of the molecule from the C-terminal residue of the HBc sequence and within about 30 residues from the C-terminus of the chimer molecule [C-terminal cysteine residue(s)],

(c) contains a sequence of at least 5 amino acid residues from HBc position 135 to the HBc C-terminus,

said chimer molecules (i) containing no more than 20 percent conservatively substituted amino acid residues in the HBc sequence, (ii) self-assembling into particles that are substantially free of binding to nucleic acids on expression in a host cell, and said particles being more stable than are particles formed from an otherwise identical HBc chimer that lacks said C-terminal cysteine residue(s) or in which a C-terminal cysteine residue present in the chimer molecule is replaced by another residue.

2. (original) The recombinant HBc chimer protein molecule according to claim 1 wherein said peptide-bonded

heterologous epitope or a heterologous linker residue for a conjugated epitope is a heterologous epitope.

3. (original) The recombinant HBc chimer protein molecule according to claim 2 wherein said heterologous epitope is a B cell epitope.

4. (original) The recombinant HBc chimer protein molecule according to claim 3 that contains a second heterologous epitope peptide-bonded to one of amino acid residues 1-4 of HBc.

5. (original) The recombinant HBc chimer protein molecule according to claim 3 wherein said B cell epitope is peptide-bonded at a position in the HBc sequence between amino acid residues 76 and 85, and at least 5 residues of the HBc sequence of positions 76 through 85 are present.

6. (original) The recombinant HBc chimer protein molecule according to claim 5 wherein the HBc sequence between amino acid residues 76 and 85 is present, but interrupted by said B cell epitope.

7. (original) The recombinant HBc chimer protein molecule according to claim 2 further including a peptide-bonded heterologous T cell epitope.

8. (original) The recombinant HBc chimer protein molecule according to claim 7 wherein said T cell epitope is peptide-bonded to the C-terminal HBc amino acid residue.

9. (original) The recombinant HBc chimer protein molecule according to claim 8 wherein said C-terminal cysteine residue(s) is present within five amino acid residues of the C-terminus of the HBc chimer protein molecule.

10-11. (cancelled)

12. (original) The recombinant HBc chimer protein molecule according to claim 1 wherein said chimer contains a heterologous linker residue for a conjugated epitope.

13. (original) The recombinant HBc chimer protein molecule according to claim 12 wherein said heterologous linker residue for a conjugated epitope is peptide-bonded at a position in the HBc sequence between amino acid residues 76 and 85, and at least 4 residues of the HBc sequence of positions 76 through 85 are present.

14. (original) The recombinant HBc chimer protein molecule according to claim 13 wherein the HBc sequence between amino acid residues 76 and 85 is present, but interrupted by said heterologous linker residue for a conjugated epitope.

15. (original) The recombinant HBc chimer protein molecule according to claim 14 that contains the HBc amino acid residue sequence of position 1 through at least position 140, plus a single cysteine residue at the C-terminus.

16. (original) The recombinant HBc chimer protein molecule according to claim 15 wherein said chimer contains the HBc amino acid residue sequence of position 1 through position 149.

17. (original) The recombinant HBc chimer protein molecule according to claim 16 wherein said heterologous linker residue for a conjugated epitope is selected from the group consisting of a lysine, aspartic acid, glutamic acid, cysteine and a tyrosine residue.

18. (currently amended) A recombinant hepatitis B virus core (HBc) protein chimer molecule with a length of about 135 to about 515 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I comprises about 71 to about 100 amino acid residues whose sequence includes at least the sequence of the residues of position 5 through position 75 of HBc and optionally includes a heterologous epitope containing up to about 30 amino acid residues peptide-bonded to one of HBc residues 1-4;

(b) Domain II comprises about 5 to about 250 amino acid residues peptide-bonded to HBc residue 75 of Domain I in which (i) zero to all residues in a sequence of HBc positions 76 through 85 are present peptide-bonded to one to about 245 amino acid residues that are heterologous to HBc and constitute a heterologous epitope or a heterologous linker residue for a conjugated epitope or (ii) ~~the sequence of HBc at positions 76 to 85 is present~~

~~free from heterologous residues, or (iii)~~ one or more of residues 76 to 85 is absent;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85 of Domain II; and

d) Domain IV comprises (i) zero through fourteen residues of a HBc amino acid residue sequence from position 136 through 149 peptide-bonded to the residue of position 135 of Domain III, (ii) one to ten cysteine residues [C-terminal cysteine residue(s)] within about 30 residues from the C-terminus of the chimer molecule, and (iii) zero to about 100 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus, with the proviso that Domain IV contain at least 6 amino acid residues including said one to ten cysteine residues of (ii),

said chimer self-assembling into particles on expression in a host cell, said particles being substantially free of binding to nucleic acids and more stable than are particles formed from an otherwise identical HBc chimer that lacks said C-terminal cysteine residue(s) or in which a C-terminal cysteine residue present in the chimer molecule is replaced by another residue, and having an amino acid residue sequence in which no more than about 10 percent of the amino acid residues are substituted in the HBc sequence of the chimer.

19. (original) The recombinant HBc chimer protein molecule according to claim 18 that contains two heterologous epitopes.

20. (original) The recombinant HBc chimer protein molecule according to claim 19 wherein said two

heterologous epitopes are present in Domains I and II, II and IV or I and IV.

21. (original) The recombinant HBc chimer protein molecule according to claim 19 wherein one of said two heterologous epitopes is a B cell epitope.

22. (original) The recombinant HBc chimer protein molecule according to claim 19 wherein one of said two heterologous epitopes is a T cell epitope.

23. (original) The recombinant HBc chimer protein molecule according to claim 19 wherein one of said two heterologous epitopes is a B cell epitope and the other is a T cell epitope.

24. (original) The recombinant HBc chimer protein molecule according to claim 18 wherein said Domain I includes a heterologous epitope peptide-bonded to one of HBc residues 1-4..

25. (original) The recombinant HBc chimer protein molecule according to claim 24 wherein said heterologous epitope of Domain II is a B cell epitope.

26. (original) The recombinant HBc chimer protein molecule according to claim 25 wherein said sequence heterologous to HBc from position 150 to the C-terminus is a T cell epitope peptide-bonded to one of HBc residues 140-149.

27. (original) The recombinant HBc chimer protein molecule according to claim 18 wherein said heterologous linker residue for a conjugated epitope or a heterologous epitope is a heterologous epitope.

28. (original) The recombinant HBc chimer protein molecule according to claim 27 wherein said heterologous epitope comprises up to about 245 amino acid residues.

29. (original) The recombinant HBc chimer protein molecule according to claim 28 wherein said heterologous epitope is a B cell epitope.

30. (original) The recombinant HBc chimer protein molecule according to claim 27 wherein said heterologous epitope contains 6 to about 50 amino acid residues.

31. (original) The recombinant HBc chimer protein molecule according to claim 27 wherein said heterologous epitope contains 20 to about 30 amino acid residues.

32. (original) The recombinant HBc chimer protein molecule according to claim 27 wherein said Domain IV comprises 1 to about 5 cysteine residues within about 30 residues from the C-terminus of the chimer molecule.

33. (original) The recombinant HBc chimer protein molecule according to claim 27 wherein the HBc

sequence between amino acid residues 76 and 85 is present, but interrupted by said heterologous epitope.

34. (cancelled)

35. (original) The recombinant HBc chimer protein molecule according to claim 18 wherein said sequence heterologous to HBc from position 150 to the C-terminus is a T cell epitope peptide-bonded to one of HBc residues 140-149.

36. (original) The recombinant HBc chimer protein molecule according to claim 18 wherein said heterologous linker residue for a conjugated epitope or a heterologous epitope is a heterologous linker residue for a conjugated epitope.

37. (original) The recombinant HBc chimer protein molecule according to claim 36 wherein said heterologous linker residue for a conjugated epitope is selected from the group consisting of a lysine, aspartic acid, glutamic acid, cysteine and a tyrosine residue.

38. (original) The recombinant HBc chimer protein molecule according to claim 37 that contains a single cysteine residue at the C-terminus of the HBc chimer protein molecule.

39-41. (cancelled)

42. (original) A recombinant hepatitis B virus core (HBc) protein chimer molecule with a length of about

175 to about 240 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I comprises about the sequence of the residues of position 1 through position 75 of HBc;

(b) Domain II comprises about 5 to about 55 amino acid residues peptide-bonded to HBc residue 75 of Domain I in which at least 4 residues in a sequence of HBc positions 76 through 85 are present peptide-bonded to 6 to about 50 amino acid residues that are heterologous to HBc and constitute a heterologous epitope;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85 of Domain II; and

d) Domain IV comprises (i) 5 through fourteen residues of a HBc amino acid residue sequence from position 136 through 149 peptide-bonded to the residue of position 135 of Domain III, (ii) a cysteine residue [C-terminal cysteine residue] within about 30 residues from the C-terminus of the chimer molecule, and (iii) zero to about 50 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus,

said chimer self-assembling into particles on expression in a host cell that exhibit a ratio of absorbance at 280 nm to 260 nm of about 1.2 to about 1.6 and are more stable than are particles formed from an otherwise identical HBc chimer molecule that lacks said C-terminal cysteine residue or in which a C-terminal cysteine residue present in the chimer molecule is replaced by another residue, and having an amino acid residue sequence in which no more than about 5 percent of the amino

acid residues are substituted in the HBc sequence of the chimer.

43. (original) The recombinant HBc chimer protein molecule according to claim 42 wherein said heterologous epitope of Domain II is a B cell epitope.

44. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein said heterologous epitope contains 15 to about 50 amino acid residues.

45. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein said heterologous epitope contains 20 to about 30 amino acid residues.

46. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein the HBc sequence between amino acid residues 76 and 85 is present, but interrupted by said heterologous epitope.

47. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein said B cell epitope is an amino acid sequence present in a pathogen selected from the group consisting of *Streptococcus pneumonia*, *Cryptosporidium parvum*, HIV, foot-and-mouth disease virus, influenza virus, *Yersinia pestis*, *Haemophilus influenzae*, *Moraxella catarrhalis*, *Porphyromonas gingivalis*, *Trypanosoma cruzi*, *Plasmodium falciparum*, *Plasmodium vivax*, *Plasmodium berghi*, *Plasmodium yoelli*, *Streptococcus sobrinus*, *Shigella flexneri*, RSV,

Plasmodium Entamoeba histolytica, *Schistosoma japonicum*,
Schistosoma mansoni, bovine inhibin and ebola virus.

48. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein said sequence heterologous to HBc from position 150 to the C-terminus is a T cell epitope peptide-bonded to one of HBc residues 140-149.

49. (original) The recombinant HBc chimer protein molecule according to claim 48 wherein said T cell epitope is from the organism against which a contemplated chimer is to be used as an immunogen.

50. (original) The recombinant HBc chimer protein molecule according to claim 43 wherein said C-terminal cysteine residue is located within about five amino acid residues of the C-terminus of the chimer protein molecule.

51. (original) An immunogenic particle comprising ~~comprised of~~ recombinant hepatitis B core (HBc) chimeric protein molecules, said chimeric protein (i) displaying one or more immunogenic epitopes at the N-terminus, HBc immunogenic loop or C-terminus, or (ii) having a heterologous linker residue for a conjugated epitope in the HBc immunogenic loop, and containing a cysteine residue at or near the C-terminus, said particle being substantially free of nucleic acid binding and exhibiting enhanced stability relative to particles comprised of otherwise identical proteins that are free of said cysteine residue.

52. (original) The immunogenic particle according to claim 51 that exhibits a 280/260 absorbance ratio of about 1.2 to about 1.7.

53. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein displays an immunogenic epitope at the N-terminus.

54. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein displays an immunogenic epitope at the C-terminus.

55. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein displays an immunogenic epitope in the immunogenic loop.

56. (currently amended) The immunogenic particle according to claim ~~4~~ 51 whose recombinant HBc chimeric protein displays a B cell immunogenic epitope.

57. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein displays a T cell immunogenic epitope.

58. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein displays separate B cell and T cell immunogenic epitopes.

59. (original) The immunogenic particle according to claim 51 whose recombinant HBc chimeric protein has a heterologous linker residue for a conjugated epitope in the HBc immunogenic loop.

60. (original) The immunogenic particle according to claim 59 wherein said heterologous linker residue for a conjugated epitope is selected from the group consisting of a lysine, aspartic acid, glutamic acid, cysteine and a tyrosine residue.

61. (original) The immunogenic particle according to claim 60 wherein said heterologous linker residue for a conjugated epitope is conjugated to a hapten.

62. (original) The immunogenic particle according to claim 61 wherein said hapten is an oligosaccharide.

63. (original) An immunogenic particle comprising ~~comprised of~~ a plurality of recombinant chimeric hepatitis B core (HBc) protein molecules;

said recombinant chimeric HBc protein molecules having a length of up to about 515 amino acid residues that

(a) contain a HBc sequence of at least about 130 of the N-terminal 150 amino acid residues of the HBc molecule that include a peptide-bonded heterologous epitope or a heterologous linker residue for a conjugated epitope present in the HBc immunodominant loop, or a sequence of at least about 135 residues of the N-terminal 150 HBc amino acid residues,

(b) contain one to ten cysteine residues toward the C-terminus of the molecule from the C-terminal residue of the HBc sequence and within about 30 residues from the C-terminus of the chimer molecule [C-terminal cysteine residue(s)],

(c) contain a sequence of at least 6 amino acid residues from HBc position 135 to the HBc C-terminus,

said chimer molecules containing no more than 10 percent conservatively substituted amino acid residues in the HBc sequence, and

said particles being substantially free of binding to nucleic acids, and being more stable than are particles formed from an otherwise identical HBc chimer that lacks said C-terminal cysteine residue(s) or in which a C-terminal cysteine residue present in the chimer molecule is replaced by another residue, and having an amino acid residue sequence in which no more than about 20 percent of the amino acid residues are substituted in the HBc sequence of the chimer.

64. (original) The immunogenic particle according to claim 63 that exhibit a ratio of absorbance at 280 nm to 260 nm of about 1.4 to about 1.6.

65. (original) The immunogenic particle according to claim 63 wherein the length of said recombinant chimeric HBc protein molecules is about 175 to about 240 amino acid residues.

66. (original) The immunogenic particle according to claim 63 wherein said peptide-bonded

heterologous epitope or a heterologous linker residue for a conjugated epitope is a heterologous epitope.

67. (original) The immunogenic particle according to claim 66 wherein said heterologous epitope is a B cell epitope.

68. (original) The immunogenic particle according to claim 63 wherein the length of said recombinant chimeric HBc protein molecules is up to about 435 amino acid residues.

69. (original) The immunogenic particle according to claim 63 that contains a second heterologous epitope peptide-bonded to one of amino acid residues 1-4 of HBc.

70. (original) The immunogenic particle according to claim 68 wherein said B cell epitope is peptide-bonded at a position in the HBc sequence between amino acid residues 76 and 85, and at least 5 residues of the HBc sequence of positions 76 through 85 are present.

71. (original) The immunogenic particle according to claim 70 wherein the HBc sequence between amino acid residues 76 and 85 is present, but interrupted by said B cell epitope.

72. (original) The immunogenic particle according to claim 68 further including a peptide-bonded heterologous T cell epitope.

73. (original) The immunogenic particle according to claim 72 wherein said T cell epitope is peptide-bonded to the C-terminal HBc amino acid residue.

74. (original) The immunogenic particle according to claim 73 wherein said C-terminal cysteine residue(s) is present within five amino acid residues of the C-terminus of the HBc chimer protein molecule.

75. (original) The immunogenic particle according to claim 63 wherein said recombinant chimeric HBc protein molecules have a length of about 135 to about 515 amino acid residues and contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I comprises about 71 to about 100 amino acid residues whose sequence includes at least the sequence of the residues of position 5 through position 75 of HBc and optionally includes a heterologous epitope containing up to about 30 amino acid residues peptide-bonded to one of HBc residues 1-4;

(b) Domain II comprises about 5 to about 250 amino acid residues peptide-bonded to HBc residue 75 of Domain I in which (i) zero to all of the residues in a sequence of HBc positions 76 through 85 are present peptide-bonded to one to about 245 amino acid residues that are heterologous to HBc and constitute a heterologous epitope or a heterologous linker residue for a conjugated epitope or (ii) the sequence of HBc at positions 76 to 85 is present free from heterologous residues;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85 of Domain II; and

d) Domain IV comprises (i) zero through fourteen residues of a HBc amino acid residue sequence from position 136 through 149 peptide-bonded to the residue of position 135 of Domain III, (ii) one to ten cysteine residues [C-terminal cysteine residue(s)] within about 30 residues from the C-terminus of the chimer molecule, and (iii) zero to about 100 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus, with the proviso that Domain IV contain at least 6 amino acid residues including said one to ten cysteine residues of (ii), said chimeric HBc protein having an amino acid residue sequence in which no more than about 10 percent of the amino acid residues are substituted in the HBc sequence.

76. (original) The immunogenic particle according to claim 75 that contains a heterologous linker residue for a conjugated epitope in Domain II and further includes a hapten linked to said heterologous linker residue.

77. (original) The immunogenic particle according to claim 76 wherein said hapten is a B cell immunogen.

78. (original) The immunogenic particle according to claim 63 wherein said recombinant chimeric HBc protein molecules have a length of about 175 to about 240 amino acid residues and contain four peptide-linked amino

acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I comprises about the sequence of the residues of position 1 through position 75 of HBc;

(b) Domain II comprises about 5 to about 55 amino acid residues peptide-bonded to HBc residue 75 of Domain I in which at least 4 residues in a sequence of HBc positions 76 through 85 are present peptide-bonded to 6 to about 50 amino acid residues that are heterologous to HBc and constitute a heterologous epitope;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85 of Domain II; and

d) Domain IV comprises (i) 5 through fourteen residues of a HBc amino acid residue sequence from position 136 through 149 peptide-bonded to the residue of position 135 of Domain III, (ii) one to about five cysteine residues [C-terminal cysteine residue] within about 30 residues from the C-terminus of the chimer molecule, and (iii) zero to about 50 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus,

said particles exhibiting a ratio of absorbance at 280 nm to 260 nm of about 1.4 to about 1.6, and said chimeric HBc protein having an amino acid residue sequence in which no more than about 5 percent of the amino acid residues are substituted in the HBc sequence.

RESPONSE UNDER 37 C.F.R. §1.116
EXPEDITED PROCEDURE
EXAMINING GROUP 1648

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant:	Ashley J. Birkett)	
)	Attorney Docket:
Serial No.:	09/931,325)	ICC-103.1 US
)	83502
Filed:	August 15, 2001)	
)	
For:	Malaria Immunogen)	Group Art Unit 1648
	and Vaccine)	
)	
Examiner:	Zachariah Lucas)	

SECOND REPLY AND AMENDMENT AFTER FINAL

Assistant Commissioner for Patents
Washington, D.C. 20231

Sir:

In reply to the Advisory Action mailed January 28, 2004 and in further reply to the Official Action mailed October 20, 2003, please amend the above-identified application as follows.

AMENDMENTS TO THE CLAIMS

1. (Currently Amended) A recombinant hepatitis B virus core (HBc) protein chimer molecule with a length of about 140 to about 310 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises the HBc sequence from position 1 through position 75 or comprises a sequence heterologous to HBc peptide-bonded to one of the first five N-terminal residues of HBc to about 85 amino acid residues whose sequence includes at least the sequence of the residues of position 5 through position 75 of HBc;

(b) Domain II comprises about 18 to about 58 amino acid residues peptide-bonded to residue 75 of which (i) a sequence of HBc is present from HBc positions 76 through 85 and (ii) a sequence of 8 to about 48 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of ~~a species of~~ the parasite *Plasmodium falciparum* that is peptide-bonded between the HBc residues of positions 78 and 79, said B cell epitope being comprised of two to about five repeats of the amino acid residue sequence Asn-Ala-Asn-Pro;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85; and

(d) Domain IV comprises a sequence of HBc from residue 136 through 140 peptide-bonded to the residue of position 135 of Domain III and (i) zero to nine residues of a HBc amino acid residue sequence from position 141 through 149, (ii) zero to three cysteine residues, (iii) fewer than

three arginine or lysine residues, or mixtures thereof adjacent to each other, and (iv) up to 100 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus, with the proviso that at least five amino acid residues are present of the amino acid residue sequence from position 136 through 149, when (a) zero cysteine residues are present and (b) fewer than about five heterologous amino acid residues are present, and

wherein no more than 10 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

2. (Original) The recombinant HBc chimer protein molecule according to claim 1 present as self-assembled particles.

3, 4, 5. (Cancelled).

6. (Original) The recombinant HBc chimer protein molecule according to claim 1 wherein Domain I consists essentially of the HBc sequence from position 1 through position 75.

7. (Original) The recombinant HBc chimer protein molecule according to claim 1 wherein Domain II independently includes zero to three peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope.

8. (Original) The recombinant HBc chimer protein molecule according to claim 1 wherein said sequence heterologous to HBc at position 150 to the C-terminus of

Domain IV comprises an amino acid residue sequence that constitutes a T cell epitope of the same species of *Plasmodium* as said B cell epitope.

9. (Currently Amended) A recombinant hepatitis B virus core (HBc) protein chimer molecule with a sequence of about 155 to about 235 amino acid residues that contains four peptide-linked domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises a sequence of residues 1 through 75 of HBc;

(b) Domain II is about 18 to about 46 residues in length of which (i) 10 residues are present in a sequence of HBc at positions 76 to 85 and (ii) a sequence of 8 to about 36 residues that constitute a repeated B cell epitope of the circumsporozoite (CS) protein of *Plasmodium falciparum* ~~or Plasmodium vivax~~ of the sequence Asn-Ala-Asn-Pro that is peptide-bonded between the residues of positions 78 and 79, said B cell epitope being comprised of two to about five repeats of an amino acid residue sequence, said Domain independently including zero to three peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope;

(c) Domain III ~~consists essentially of~~ comprises the HBc sequence from position 86 through position 135; and

(d) Domain IV comprises a sequence of HBc from residue 136⁷ through 140 peptide-bonded to the residue of position 135 of Domain III and (i) zero to nine residues of a HBc amino acid residue sequence from position 141 through 149, (ii) ~~zero~~ one to three cysteine residues, (iii) fewer than three arginine or lysine residues, or mixtures thereof

adjacent to each other, and (iv) up to 50 amino acid residues in a sequence that constitutes a T cell epitope of Plasmodium falciparum ~~the same species of Plasmodium as said B cell epitope~~ peptide-bonded to the final HBc amino acid residue present in the chimer, and

wherein no more than 10 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

10. (Original) The recombinant HBc chimer protein molecule according to claim 9 wherein Domain IV comprises one amino acid residue to a sequence of about nine amino acid residues of the HBc sequence from residue position 141 through about position 149 peptide-bonded to residue 140.

11. (Original) The recombinant HBc chimer protein molecule according to claim 10 wherein Domain IV consists essentially of a sequence of nine amino acid residues of the HBc sequence from residue position 141 through position 149 peptide-bonded to residue 140.

12, 13. (Cancelled).

14. (Currently Amended) The recombinant HBc chimer protein molecule according to claim 9 ~~13~~ wherein the repeated sequence of said B cell epitope of Domain II is repeated three or four times.

15. (Original) The recombinant HBc chimer protein molecule according to claim 14 wherein the repeated

sequences are peptide-bonded to each other without interruption.

16. (Currently Amended) The recombinant HBc chimer protein molecule according to claim 15 wherein said B cell epitope includes a second CS protein sequence from ~~the same~~ *Plasmodium falciparum* ~~species~~ that is peptide-bonded to said repeated sequence.

17. (Original) The recombinant HBc chimer protein molecule according to claim 16 wherein said second CS protein sequence is Asn-Val-Asp-Pro.

18. (Cancelled).

19. (Original) The recombinant HBc chimer protein molecule according to claim 17 wherein said second CS protein sequence is peptide-bonded at the amino-terminus of said repeated sequence.

20. (Original) The recombinant HBc chimer protein molecule according to claim 16 wherein said second CS protein sequence is SEQ ID NO:126 (Asn-Ala-Asn-Pro-Asn-Val-Asp-Pro).

21. (Cancelled).

22. (Original) The recombinant HBc chimer protein molecule according to claim 20 wherein said second CS protein sequence is peptide-bonded at the amino-terminus of said repeated sequence.

23. (Currently Amended) The recombinant HBc chimer protein molecule according to claim 10 wherein said one to three cysteine residues are present in Domain IV within about 30 residues of the carboxy-terminus of the chimeric molecule.

24. (Original) The recombinant HBc chimer protein molecule according to claim 23 wherein said one to three cysteine residues are present in said T cell epitope.

25. (Currently Amended) The recombinant HBc chimer protein molecule according to claim 24 wherein said T cell epitope is present and has the sequence of SEQ ID NO: 148 or 24 25 (EYLNKIQNSLSTEWSPCSVT OR GIEYLNKIQNSLSTEWSPCSVT-YLDKVRATVGTETWTPCSVT).

26. (Original) The recombinant HBc chimer protein molecule according to claim 9 present as self-assembled particles.

27. (Currently Amended) Particles comprised of recombinant hepatitis B virus core (HBc) protein chimer molecules, said molecules having a sequence of about 155 to about 235 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises the HBc sequence from position 1 through position 75 or comprises a sequence heterologous to HBc peptide-bonded to one of the first five N-terminal residues of HBc to about 85 amino acid residues whose sequence includes at least the

sequence of the residues of position 5 through position 75 of HbC;

(b) Domain II comprises about 18 to about 58 amino acid residues peptide-bonded to residue 75 of which (i) a sequence of HbC is present from HbC positions 76 through 85 and (ii) a sequence of 8 to about 48 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of ~~a species of the parasite~~ *Plasmodium falciparum* that is peptide-bonded between the HbC residues of positions 78 and 79, said B cell epitope being comprised of two to about five repeats of an amino acid residue sequence Asn-Ala-Asn-Pro;

(c) Domain III is an HbC sequence from position 86 through position 135 peptide-bonded to residue 85; and

(d) Domain IV comprises a sequence of HbC from residue 136 through 140 peptide-bonded to the residue of position 135 of Domain III, and (i) zero to nine residues of a HbC amino acid residue sequence from position 141 through 149, (ii) zero to three cysteine residues, ~~(iii) zero to three cysteine residues,~~ (iii) fewer than three arginine or lysine residues, or mixtures thereof adjacent to each other, and (iv) up to 50 amino acid residues in a sequence heterologous to HbC from position 150 to the C-terminus ~~with the proviso that at least five amino acid residues are present of the HbC amino acid residue sequence from position 136 through 149 when (a) zero cysteine residues are present and (b) fewer than about five heterologous amino acid residues are present,~~ and

wherein no more than 10 percent of the HbC amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

28. (Original) The particles according to claim 27 whose HBc chimer protein molecules have a sequence length of about 165 to about 210 amino acid residues.

29, 30, 31. (Cancelled).

32. (Original) The particles according to claim 27 wherein Domain I consists essentially of the HBc sequence from position 1 through position 75.

33. (Original) The particles according to claim 27 wherein Domain II independently includes zero to three peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope.

34. (Currently Amended) The particles according to claim 27 wherein said sequence heterologous to HBc at position 150 to the C-terminus of Domain IV comprises an amino acid residue sequence that constitutes a T cell epitope of Plasmodium falciparum ~~the same species of Plasmodium as said B cell epitope.~~

35. (Currently Amended) Particles comprised of recombinant hepatitis B virus core (HBc) protein chimer molecules, said molecules having a sequence of about 165 to about 210 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises a sequence of residues 1 through position 75 of HBc;

(b) Domain II comprises about 18 to about 46 amino acid residues peptide-bonded to residue 75 of which (i) 10 residues are present in a sequence of HBc from position 76 through 85 and (ii) a sequence of 8 to about 36 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of *Plasmodium falciparum* or ~~*Plasmodium vivax*~~ that is peptide-bonded between the residues of HBc positions 78 and 79, said B cell epitope being comprised of two to about five repeats of the an amino acid residue sequence Asn-Ala-Asn-Pro, said Domain independently including zero to two peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope;

(c) Domain III ~~consists essentially of the~~ is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85; and

(d) Domain IV comprises the HBc sequence of residues 136 through 140 peptide-bonded to the residue of position 135 of Domain III and (i) zero to nine residues of a HBc amino acid residue sequence from position 140 through 149 peptide-bonded to the residue of position 140, (ii) zero one to three cysteine residues, (iii) fewer than three arginine or lysine residues, or mixtures thereof adjacent to each other, and (iv) up to 25 amino acid residues in a sequence that constitutes a T cell epitope of the same species of *Plasmodium* as said B cell epitope, said T cell epitope sequence being peptide-bonded to the final HBc amino acid residue present in a chimer molecule or a cysteine residue, and

wherein no more than 10 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

36. (Currently Amended) The particles according to claim 35 wherein Domain IV comprises one to a sequence of nine amino acid residues of the HBc sequence from residue position 141 through position 149 linked between residue 140 of said Domain III sequence and a *Plasmodium falciparum* or ~~*Plasmodium vivax*~~ T cell epitope.

37. (Original) The particles according to claim 36 wherein the nine amino acid residues of the HBc sequence from residue position 141 through position 149 are present.

38, 39. (Cancelled).

40. (Original) The particles according to claim 39 wherein the repeated sequence of said B cell epitope of Domain II is repeated three or four times.

41. (Original) The particles according to claim 40 wherein the repeated sequences are peptide-bonded to each other without interruption.

42. (Currently Amended) The particles according to claim 41 wherein said B cell epitope includes a second CS protein sequence from *Plasmodium falciparum* ~~the same *Plasmodium* species~~ that is peptide-bonded to said repeated sequence.

43. (Original) The particles according to claim 42 wherein said second CS protein sequence is Asn-Val-Asp-Pro.

44. (Original) The particles according to claim 43 wherein said second CS protein sequence is peptide-bonded at the carboxy-terminus of said repeated sequence.

45. (Original) The particles according to claim 43 wherein said second CS protein sequence is peptide-bonded at the amino-terminus of said repeated sequence.

46. (Original) The particles according to claim 42 wherein said second CS protein sequence is SEQ ID NO:126 (Asn-Ala-Asn-Pro-Asn-Val-Asp-Pro).

47. (Original) The particles according to claim 46 wherein said second CS protein sequence is peptide-bonded at the carboxy-terminus of said repeated sequence.

48. (Original) The particles according to claim 46 wherein said second CS protein sequence is peptide-bonded at the amino-terminus of said repeated sequence.

49. (Original) The particles according to claim 35 wherein said B cell epitope of *Plasmodium falciparum* has an amino acid residue sequence selected from the group consisting of SEQ ID NOS:1-14.

50. (Cancelled).

51. (Original) The particles according to claim 35 wherein said T cell epitope of *Plasmodium falciparum* is present and has the amino acid residue sequence of SEQ ID NO:24.

52. (Cancelled).

53. (Currently Amended) The particles according to claim 36 ~~further including wherein said~~ one to three cysteine residues in the Domain IV sequence are present within about 30 residues of the carboxy-terminus of the chimeric molecule.

54. (Original) The particles according to claim 53 having one cysteine residue in the Domain IV sequence.

55. (Cancelled).

56. (Currently Amended) Particles comprised of recombinant hepatitis B virus core (HBc) protein chimer molecules, said molecules having a sequence of about 165 to about 210 amino acid residues that contain four peptide-linked domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises a sequence of residues 1 through position 75 of HBc;

(b) Domain II comprises about 18 to about 46 amino acid residues peptide-bonded to residue 75 of which (i) 10 residues are present in a sequence of HBc from position 76 through 85 and (ii) a sequence that constitutes a B cell epitope of the circumsporozoite (CS) protein of *Plasmodium falciparum* ~~or Plasmodium vivax~~ is peptide-bonded between the residues of HBc positions 78 and 79, said B cell epitope being selected from the group consisting of SEQ ID NOs: 1-14 ~~1-21~~, said Domain II including two peptide-bonded residues on either side of said B cell

epitope that are other than those of HBc or said B cell epitope;

(c) Domain III ~~consists essentially of the~~ is an HBc sequence from position 86 through position 135 peptide bonded to residue 85; and

(d) Domain IV comprises the sequence of HBc residues 136-140 peptide-bonded to residue 135 plus one to nine residues of a HBc amino acid residue sequence from position 141 through 149 peptide-bonded to the residue of position 140 and also peptide-bonded to a *Plasmodium falciparum* ~~or Plasmodium vivax~~ T cell epitope of a sequence of up to about 25 amino acid residues that includes a cysteine residue, and

wherein no more than 10 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

57. (Currently Amended) The particles according to claim 56 wherein Domain IV comprises nine amino acid residues of the HBc sequence from residue position 141 through position 149 bonded between said residue 140 and said *Plasmodium falciparum* ~~or Plasmodium vivax~~ T cell epitope.

58. (Currently Amended) The particles according to claim 57 wherein ~~said B cell epitope is of the CS protein of Plasmodium falciparum that is selected from the group consisting of SEQ ID NOs:1-14 and said Plasmodium falciparum T cell epitope has the amino acid sequence of~~ SEQ ID NO:24.

59. Cancelled.

60. (Currently Amended) A vaccine or inoculum comprising an immunogenic effective amount immunogenic particles dissolved or dispersed in a pharmaceutically acceptable diluent, wherein said immunogenic particles are comprised of a plurality of recombinant chimeric hepatitis B core (HBc) protein molecules having a length of about 140 to about 310 amino acid residues that contain four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises the HBc sequence from position 1 through position 75 or comprises a sequence heterologous to HBc peptide-bonded to one of the first five N-terminal residues of HBc to about 85 amino acid residues whose sequence includes at least the sequence of the residues of position 5 through position 75 of HBc;

(b) Domain II comprises about 18 to about 58 amino acid residues peptide-bonded to residue 75 of which (i) a sequence of HBc is present from HBc positions 76 through 85 and (ii) a sequence of 8 to about 48 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of ~~a species of the parasite~~ *Plasmodium falciparum* that is peptide-bonded between the HBc residues of positions 78 and 79, said B cell epitope being comprised of two to about five repeats of the amino acid residue sequence Asn-Ala-Asn-Pro;

(c) Domain III is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85; and

(d) Domain IV comprises a sequence of HBc from residue 136 through 140 peptide-bonded to the residue of

position 135 of Domain III and (i) zero to nine residues of a HBc amino acid residue sequence from position 141 through 149, (ii) zero to three cysteine residues, ~~(ii) zero to three cysteine residues,~~ (iii) fewer than three arginine or lysine residues, or mixtures thereof adjacent to each other, and (iv) up to 100 amino acid residues in a sequence heterologous to HBc from position 150 to the C-terminus, ~~with the proviso that at least five amino acid residues are present of the amino acid residue sequence from position 136 through 149, when (a) zero cysteine residues are present and (b) fewer than about five heterologous amino acid residues are present, and~~

wherein no more than 10 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

61. (Currently Amended) The vaccine or inoculum according to claim 60 wherein said immunogenic particles are those having a sequence of about 165 to about 210 amino acid residues that contains four peptide-linked amino acid residue sequence domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises a sequence of residues 1 through position 75 of HBc;

(b) Domain II comprises about 18 to about 46 amino acid residues peptide-bonded to residue 75 of which (i) 10 residues are present in a sequence of HBc from position 76 through 85 and (ii) a sequence of 8 to about 36 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of *Plasmodium falciparum* ~~or Plasmodium vivax~~ that is peptide-bonded between the residues of HBc positions 78 and 79, said B cell epitope

being comprised of two to about five repeats of an the amino acid residue sequence Asn-Ala-Asn-Pro, said Domain independently including zero to two peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope;

(c) Domain III ~~consists essentially of the~~ is an HBc sequence from position 86 through position 135 peptide-bonded to residue 85; and

(d) Domain IV comprises the HBc sequence of residues 136 through 140 peptide-bonded to the residue of position 135 of Domain III and (i) nine residues of a HBc amino acid residue sequence from position 141 through 149 peptide-bonded to the residue of position 140, (ii) one to three cysteine residues, (iii) fewer than three arginine or lysine residues, or mixtures thereof adjacent to each other, and (iv) a *Plasmodium falciparum* ~~or Plasmodium vivax~~ T cell epitope, said T cell epitope sequence being peptide-bonded to the final HBc amino acid residue present in a chimer molecule or a cysteine residue, and

wherein no more than 5 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

62. (Previously Presented) The vaccine or inoculum according to claim 60 wherein Domain II comprises about 18 to about 46 amino acid residues peptide-bonded to residue 75 of which (i) 10 residues are present in a sequence of HBc from position 76 through 85 and (ii) a sequence of 8 to about 36 residues that constitute a B cell epitope of the circumsporozoite (CS) protein of *Plasmodium falciparum* that is peptide-bonded between the residues of HBc positions 78 and 79, said B cell epitope being

comprised of three or four repeats of an amino acid residue sequence Asn-Ala-Asn-Pro, said Domain independently including zero to two peptide-bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope.

63. (Currently Amended) The vaccine or inoculum according to claim 62 wherein the repeated sequences are peptide-bonded to each other without interruption and wherein said B cell epitope includes a second CS protein sequence from Plasmodium falciparum ~~the same Plasmodium species~~ that is peptide-bonded to said repeated sequence.

64. (Currently Amended) The vaccine or inoculum according to claim 60 wherein said immunogenic particles are those comprised of recombinant hepatitis B virus core (HBc) protein chimer molecules, said molecules having a sequence of about 165 to about 210 amino acid residues that contain four peptide-linked domains from the N-terminus that are denominated Domains I, II, III and IV, wherein

(a) Domain I ~~consists essentially of~~ comprises a sequence of residues 1 through position 75 of HBc;

(b) Domain II comprises about 18 to about 46 amino acid residues peptide-bonded to residue 75 of which (i) 10 residues are present in a sequence of HBc from position 76 through 85 and (ii) a sequence that constitutes a B cell epitope of the circumsporozoite (CS) protein of Plasmodium falciparum ~~ex-Plasmodium vivax~~ is peptide-bonded between the residues of HBc positions 78 and 79, said B cell epitope being selected from the group consisting of SEQ ID NOs: 1-141-21, said Domain II including two peptide-

bonded residues on either side of said B cell epitope that are other than those of HBc or said B cell epitope;

(c) Domain III ~~consists essentially of the~~ is an HBc sequence from position 86 through position 135 peptide bonded to residue 85; and

(d) Domain IV comprises the sequence of HBc residues 136-140 peptide-bonded to residue 135 plus nine residues of a HBc amino acid residue sequence from position 141 through 149 peptide-bonded to the residue of position 140 and also peptide-bonded to a *Plasmodium falciparum* T cell epitope of a sequence of up to about 25 amino acid residues that includes a cysteine residue, and

wherein no more than 5 percent of the HBc amino acid residues are substituted as compared to SEQ ID NO:170 from position 1 through 149.

65. (Currently Amended) The vaccine or inoculum according to claim 64 wherein said immunogenic particles are those wherein ~~said B cell epitope is of the CS protein of Plasmodium falciparum that is selected from the group consisting of SEQ ID NOs:1-14 and said Plasmodium falciparum T cell epitope has the amino acid sequence of~~ SEQ ID NO:24.

66. (Cancelled).

67. (Original) The vaccine or inoculum according to claim 60 that is adapted for parenteral administration.

68-75. (Cancelled).

Department of Health and Human Services Public Health Service Small Business Innovation Research Program Phase I Grant Application <i>Follow instructions carefully.</i>	Leave blank — for PHS use only.		
	Type	Activity	Number
	Review Group		Formerly
	Council Board (Month, year)		Date Received

1. TITLE OF APPLICATION (Do not exceed 56 typewriter spaces)

Malaria Vaccine

2. SOLICITATION NO. PHS 97-2

3. PRINCIPAL INVESTIGATOR

☐ New Investigator

3a. NAME (Last, first, middle)

Birkett, Ashley James

3b. DEGREE(S)

B.S. Ph.D.

3c. SOCIAL SECURITY NO.

Provide on Personal Data Page

3d. POSITION TITLE

Director of Biochemistry

3e. MAILING ADDRESS (Street, city, state, zip code)

3347 Industrial Court, Suite A
San Diego, CA 92121

3f. TELEPHONE AND FAX (Area code, number, and extension)

TEL: (619) 793-2661

FAX: (619) 793-2666

BITNET/INTERNET Address:

abirkett@ix.netcom.com

4. HUMAN SUBJECTS

4a. If "yes," Exemption no.

or ☐

IRB approval date

☐ Full IRB or
Expedited
Review4b. Assurance of
compliance no.5. VERTEBRATE
ANIMALS5a. If "yes,"
IACUC
approval
date5b. Animal welfare
assurance no.☒ NO☐ YES☐ NO☒ YES

7-14-94

A-2430-01

6. DATES OF PROJECT PERIOD

From: 7/1/98

Through: 12/31/98

7. COSTS REQUESTED

7a. Direct Costs

\$ 80,000

7b. Total Costs

\$100,000

8. PERFORMANCE SITES (Organizations and addresses)

ICC/Synthetic Genetics
3347 Industrial Court, Suite A
San Diego, CA 921219. APPLICANT ORGANIZATION (Name and address of applicant
small business concern)Ashley J. Birkett, Ph.D.
ICC/Synthetic Genetics
3347 Industrial Court, Suite A
San Diego, CA 92121

10. ENTITY IDENTIFICATION NUMBER

1-95-446-7306-A1

Congressional District

41st

11. SMALL BUSINESS CERTIFICATION

☒ Small Business Concern☐ Women-owned☐ Socially and Economically Disadvantaged12. NOTICE OF PROPRIETARY INFORMATION: The information identified
on all pages

of this application constitutes trade secrets or information that is commercial or financial and confidential or privileged. It is furnished to the Government in confidence with the understanding that such information shall be used or disclosed only for evaluation of this application, provided that, if a grant is awarded as a result of or in connection with the submission of this application, the Government shall have the right to use or disclose the information herein to the extent provided by law. This restriction does not limit the Government's right to use the information if it is obtained without restriction from another source.

13. DISCLOSURE PERMISSION STATEMENT: If this application does not result in an award, is the Government permitted to disclose the title only of your proposed project, and the name, address, and telephone number of the official signing for the applicant organization, to organizations that may be interested in contacting you for further information or possible investment? ☐ YES ☒ NO

14. OFFICIAL SIGNING FOR APPLICANT ORGANIZATION

Name: George B. Thornton

Title: President/CEO

Address: ICC/Synthetic Genetics
3347 Industrial Court, Suite A
San Diego, CA 92121

Telephone: (619) 793-2661

FAX: (619) 793-2666

BITNET/INTERNET Address:

bthorn@ix.netcom.com

15. PRINCIPAL INVESTIGATOR ASSURANCE: I certify that the statements herein are true, complete, and accurate to the best of my knowledge. I am aware that any false, fictitious, or fraudulent statements or claims may subject me to criminal, civil, or administrative penalties. I agree to accept responsibility for the scientific conduct of the project and to provide the required progress reports if a grant is awarded as a result of this application.

SIGNATURE OF PERSON NAMED IN 3a
(In ink. "Per" signature not acceptable.)

DATE

12/11/97

16. APPLICANT ORGANIZATION CERTIFICATION AND ACCEPTANCE: I certify that the statements herein are true, complete, and accurate to the best of my knowledge, and accept the obligation to comply with Public Health Service terms and conditions if a grant is awarded as a result of this application. I am aware that any false, fictitious, or fraudulent statements or claims may subject me to criminal, civil, or administrative penalties.

SIGNATURE OF PERSON NAMED IN 14
(In ink. "Per" signature not acceptable.)

DATE

12/11/97

Abstract of Research Plan

NAME, ADDRESS, AND TELEPHONE NUMBER OF APPLICANT ORGANIZATION

ICC/Synthetic Genetics
3347 Industrial Court, Suite A
San Diego, CA 92121
Telephone: (619) 793-2661

YEAR FIRM FOUNDED

1994

NO. OF EMPLOYEES (include all affiliates)

15

TITLE OF APPLICATION

KEY PERSONNEL ENGAGED ON PROJECT

NAME	ORGANIZATION	ROLE ON PROJECT
Ashley J. Birkett, Ph.D.	ICC/SG	Principal Investigator

ABSTRACT OF RESEARCH PLAN: State the application's broad, long-term objectives and specific aims, making reference to the health-relatedness of the project. Describe concisely the research design and methods for achieving these goals and discuss the potential of the research for technological innovation. Avoid summaries of past accomplishments and the use of the first person. This abstract is meant to serve as a succinct and accurate description of the proposed work when separated from the application. If the application is funded, this description, as is, will become public information. *Therefore, do not include proprietary or confidential information.* DO NOT EXCEED 200 WORDS.

Pre-erythrocytic stage malaria vaccines aim to block hepatocyte entry by sporozoites and/or release of merozoites into the blood stream, thereby circumventing the disease process and rendering the host non-infectious. The primary target of this class of vaccines is the circumsporozoite protein (CS), which is a major constituent of the sporozoite coat when it enters the host. Pre-erythrocytic immunity has been demonstrated by immunization with the CS-NANP repeat epitope, passive transfer of antibodies specific for anti-NANP repeat epitope, and immunization with irradiated sporozoites.

In animal models the immunodominant CS-repeat epitope from either *P.bergei* or *P.yoelli*, displayed on hepatitis B core particles, successfully protected >90% of animals against infection. Preliminary studies of HBc particles engineered to deliver the CS-repeat from the human parasite *P.falciparum* are promising, but in need of optimization.

The focus of this work is to optimize the immunogenicity of the *P.falciparum* particle to attain the high immunogenicity and absence of genetic restriction observed with *P.bergei* and *P.yoelli* particles. This will be achieved by optimizing the presentation of the NANP repeat epitope at the surface of HBc and incorporating a universal malaria-specific T cell epitope. Once identified, this vaccine candidate will be the subject of clinical testing during phase II.

Provide key words (8 maximum) to identify the research or technology.

malaria, plasmodium falciparum, circumsporozoite, vaccine

Provide a brief summary of the potential commercial applications of the research.

Malaria is by far the world's most important tropical parasitic disease, and kills more people than any other communicable disease, with the exception of tuberculosis. Malaria is a public health problem in more than 90 countries, inhabited by a total of 2.4 billion people – 40% of the world's population. Mortality due to malaria is estimated to be in the range of 1.5 to 2.7 million deaths each year, accounting for one person every 12 seconds. There are 7 million travelers from the U.S. each year to endemic areas.

Budget for Phase I—Direct Costs Only

FROM

6/1/98

TO

11/30/98

PERSONNEL (Applicant organization only)

NAME	Role on Project	Type Appt. (months)	% Effort on Project	Institutional Base Salary	DOLLAR AMOUNT REQUESTED (omit cents)		
					Salary Requested	Fringe Benefits	TOTALS
Ashley J. Birkett	P.I.	12	100	60,000	30,000	4,500	34,500
Katie Lyons	Tech.	12	100	24,000	12,000	1,800	13,800
Romeo Veniegas	Tech.	12	50	24,000	6,000	900	6,900
SUBTOTALS →					48,000	7,200	55,200

CONSULTANT COSTS

David R. Milich, Ph.D. (30 hrs @ \$50/hr)
 Ruth S. Nussenzweig, Ph.D. (30 hrs @ \$50/hr)

3,000

EQUIPMENT (Itemize)

None

0

SUPPLIES (Itemize by category)

Chemicals and Reagents \$7,500
 Plasticware and Glassware \$2,500

10,000

TRAVEL

None

0

PATIENT CARE COSTS

Inpatient None

Outpatient None

0

CONTRACTUAL COSTS

0

OTHER EXPENSES (Itemize by category)

Animal Purchase/Service (14, 4-Mice Protocols @ \$500/per)
 Oligonucleotide Synthesis/Purification (\$800)
 DNA Sequencing (\$600)
 Electron Microscopy (8 @ \$425/per ABI) (\$3,400)

11,800

TOTAL DIRECT COSTS (Also enter on Face Page, Item 7a) →

\$ 80,000

FIXED FEE REQUESTED

\$ 0

OTHER SUPPORT (see instructions)



NO



YES

Budget Justification

Using continuation pages if necessary, describe the specific functions of the personnel and consultants. Read the instructions and justify costs accordingly.

Personnel/Consultant Costs

Ashley J. Birkett, Ph.D.	Oversee project experimental design, experimentation and data analysis; interact with scientific advisory board members; purification and physical/biochemical analysis of particles; prepare Phase I report and Phase II proposal.
Kate Lynne Lyons, B.S.	Cloning and purification of particles, stability studies, antisera testing.
Romeo Veneigas, B.S.	General laboratory support (reagent preparation, dishwashing, ordering), fermentation.
David R. Milich, Ph.D.	Consult on immunological aspects of projects, including experimental design and data analysis of genetic restriction studies.
Ruth S. Nussenzweig, Ph.D.	Assist in data analysis and selection of the lead candidate. The expertise she has gained from her lifelong studies on malaria and her involvement in previous clinical trials of multiple malaria vaccine candidates will be invaluable to this proposal.

Reagent Costs

Chemicals and Reagents: Cloning reagents (agarose, restriction/modifying enzymes, gel extraction kit, plasmid purification kits); Fermentation media (casein, yeast extract, glucose, ampicillin, agar); Purification reagents (ammonium sulfate, Sepharose CL-4B, HA); Protein Analysis reagents (acrylamide, membranes, SDS, Polaroid film); ELISA reagents (Anti-mouse peroxidase conjugates, TM Blue substrate); Salts (Tris, NaCl, sodium phosphate, etc.)

Plasticware and Glassware: Disposable glass tubes, Eppendorf tubes, pipette tips, serological pipettes, cuvettes, microtitre plates, culture dishes, centrifuge bottles and tubes, chromatography columns, etc.

Animal Services: 14 x 4 Mice Protocols

Resources

FACILITIES: Specify the facilities to be used for the conduct of the proposed research. Indicate their capacities, pertinent capabilities, relative proximity, and extent of availability to the project. Include laboratory, clinical, animal, computer, and office facilities at the applicant small business concern and any other performance site listed on the FACE PAGE. Identify support services such as secretarial, machine shop, electronics shop, and the extent to which they will be available to the project. Use continuation page(s) if necessary.

ICC/SG's 7500 sq.ft. is located in the Sorrento Valley district of San Diego, proximal to UCSD and The Scripps Research Institute, a popular location for Biotech companies. The facility includes 5000 sq.ft. of wet lab space and 2500 sq.ft. of office space with full-time secretarial, marketing and accounting services available for this project. The company has full internet access in addition to access to scientific literature via the comprehensive bio-medical library at UCSD. The laboratories are fully equipped and engaged in ongoing research funded by NIH, NIEHS, product sales from its subsidiary Synthetic Genetics, and private investment. No animal facilities are available on-site; animal studies are routinely contracted out to commercial establishments.

MAJOR EQUIPMENT: List the most important equipment items already available for this project, noting the location and pertinent capabilities of each.

Floor and benchtop centrifuges, microfuges, UV/Vis spectrophotometers, analytic and semi-preparative HPLCs with autosamplers and integrators, kinetic ELISA plate reader, chart recorders, fraction collectors, peristaltic pumps, in-line UV/Vis monitors, computer controlled purification systems (Bio-Rad and PerSeptive BioCad), French press, fume hood, various shakers, waterbaths, PCR thermocyclers, refrigerated chromatography cabinet, -20°C and -70°C degree freezers, agarose and PAGE electrophoresis units, power supplies, darkroom with UV box, gel documentation system, computers, printers, scanner, photocopiers.

A. SPECIFIC AIMS

The focus of this work is to optimize the B cell and T cell immunogenicity of the *P.falciparum* particle such that it has the high immunogenicity and absence of genetic restriction achieved with *P.bergei* and *P.yoelli*, which were protective to levels of 90-100% in rodents [1, 2]. This will be achieved by optimizing the presentation of the NANP repeat epitope at the surface of the HBc carrier using an improved presentation technique, and incorporating a recently identified universal malaria-specific T cell epitope.

1. Incorporation of the Malaria Specific T cell epitope

HBc is highly effective in enhancing antibody responses to 'carried' epitopes in a manner which shows little or no genetic restriction, therefore ensuring universal antibody priming. This is achieved through a combination of the architecture of the HBc particle, efficient antigen presentation of HBc by B cells and potent T cell recognition of HBc. To ensure the priming of malaria specific T_H cells as well as malaria-specific antibody producing B cells, we intend to incorporate a recently identified universal malaria-specific T cell epitope (Pf Th/Tc) in our vaccine [3, 4].

The importance of including a universal T_H epitope derived from the malaria parasite in a malaria vaccine are several fold. First, the inclusion of an additional T_H epitope will potentially help to increase antibody responses to the NANP epitope in individuals where recognition of the HBc T_H epitope may be limiting. Secondly, the priming of malaria specific T_H cells ensures an anamnestic response to *P.falciparum* such that, should a vaccine recipient be exposed to malaria, a more rapid and stronger anti-malaria response will be activated due to previous priming of malaria specific T_H cells. Thirdly, vaccinees living in malaria endemic regions will experience natural 'boosting' every time they are exposed to the parasite because their immune systems have been primed at both the B and T_H cell level. This effect is similar to clinical boosting by re-vaccination – a phenomenon which is particularly attractive for third-world prophylaxis in areas where malaria is endemic.

The malaria-specific T cell site we have selected for inclusion in our vaccine is a universal form of the T cell epitope CS 326-345, recently identified from T cell clones of volunteers who became protected against malaria infection following immunization with irradiated sporozoites [3, 4]. The data presented shows that this epitope appears to bind all human MHC class II molecules and is therefore 'universal' in nature, suggesting that it will prime malaria specific T_H cells in essentially all vaccine recipients. Preliminary reports also indicate that this epitope primes class I restricted T cells. This universal T_H/T_c epitope will be genetically fused to the HBc/NANP particles and tested for its functionality using a variety of *in vitro* and *in vivo* assays.

2. Optimizing Anti-NANP Immunogenicity

The CS protein of *P.falciparum* contains approximately 40 NANP repeats; however, the immunodominant CS epitope is contained within just 3 repeats [5]. To date, our studies have focused on HBc particles bearing 4 repeats at the immunodominant loop region of HBc [1, 2]. We have recently shown that variation of the precise location of the epitope in the immunodominant loop can increase immunogenicity (unpublished observation). Since HBc is a 180 subunit particle, optimum immunogenicity must be determined for this particular presentation system as the conformation of the inserted epitope will vary depending upon the number of repeats presented, which may in turn translate to variable immunogenicity. To address this issue we will investigate the effects of displaying either 3, 6, 9 or 12 NANP repeats at the surface of HBc particles bearing the universal T_H epitope [3]. These particles will be used to immunize mice from multiple genetic backgrounds to investigate immunogenicity and genetic restriction of the anti-NANP immune response.

3. Stability Studies

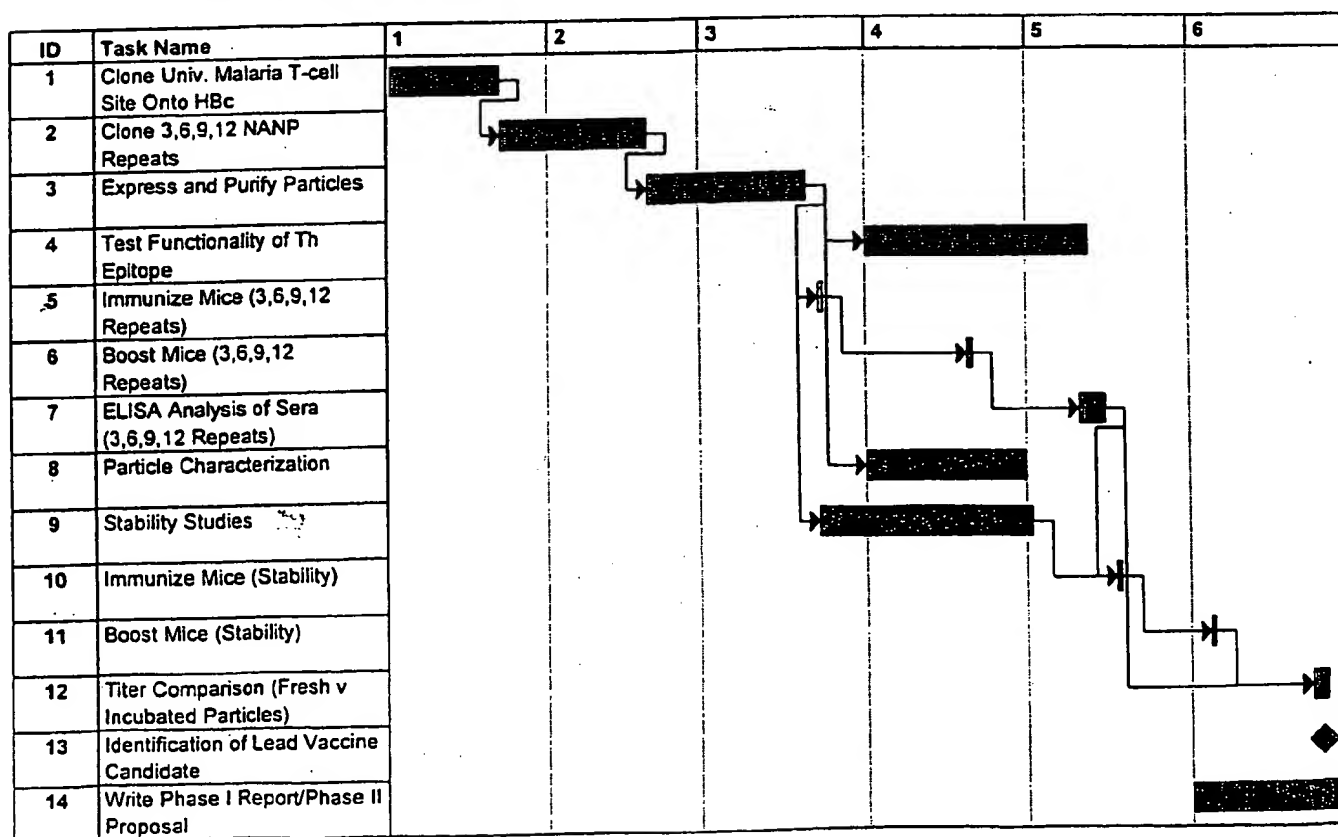
Two critical concerns of an effective malaria vaccine projected for use in the third world are stability and cost. Cost considerations are discussed later. Initial studies of HBc particles show them to be extremely thermostable, an important consideration given the potential use of this vaccine in developing countries. To address stability of the proposed constructs, accelerated stability studies of the four HBc/NANP particles described in 'Specific Aims 2' will be conducted at 37°C. Particles will be examined by SDS-PAGE to detect proteolysis, electron microscopy to examine particle integrity, ELISA for particle antigenicity, and immunogenicity using B10.S mice. The properties of 'incubated particles' will be critically compared with those stored at -70°C degrees.

Criteria for the Successful Completion of Phase I

The overall aim of the phase I funding period is to optimize the stability and immunogenicity of the NANP/Universal T cell epitope-containing particle. The criteria for the identification and subsequent clinical development (SBIR Phase II) of a candidate particle will be the following:

1. Immunogenicity levels in mice comparable to those achieved in the *P.bergeii* rodent model across multiple genetic backgrounds.
2. Yields of purified hybrid particles which exceed 50 mg/L, which will therefore minimize production costs.
3. Acceptable particle stability as determined by accelerated stability studies conducted at 37°C. Our goal is to develop a particle which is stable in solution for several months at ambient temperature.
4. Minimize the requirement for adjuvants beyond those currently approved (i.e. alum).

Timeline for the Completion of Phase I



B. SIGNIFICANCE

Malaria

Malaria is by far the world's most important tropical parasitic disease, and kills more people than any other communicable disease, with the exception of tuberculosis. The causative agents in humans are four species of *Plasmodium* protozoa: *P.falciparum*, *P.vivax*, *P.ovale* and *P.malariae*. Of these, *P.falciparum* accounts for the majority of infections and is the most lethal.

- Malaria is a public health problem today in more than 90 countries, inhabited by a total of 2.4 billion people – 40% of the world's population.
- Worldwide prevalence of the disease is estimated to be on the order of 300-500 million clinical cases each year.
- More than 90% of all malaria cases are in sub-Saharan Africa. Two-thirds of the remainder are concentrated in six countries: India, Brazil, Sri Lanka, Viet Nam, Columbia and Solomon Islands, in decreasing order of prevalence.
- Mortality due to malaria is estimated to be in the range of 1.5 to 2.7 million deaths each year, accounting for one person every 12 seconds. The vast majority of deaths occur among young children in Africa, especially in remote rural areas with poor access to health services.
- The death rate of children below the age of 5 years is high but older children and adults generally survive, although they may have bouts of illness and may continually carry the organism.
- There are 7 million travelers from the U.S. each year to endemic areas. One source estimates that there are as many as 20 million arrivals from the U.S. into countries, other than Africa, where malaria is prevalent (World Tourism Organization).
- The U.S. reports about 1,000 cases of malaria each year. About 10,000 cases are reported each year in Europe, alone.

(Data from , WHO Factsheet #94, December 1996 and [44])



Figure 1: Malaria is a public health problem today in more than 90 countries, inhabited by a total of 2.4 billion people – 40% of the world's population.

Malaria infection begins when a person is bitten by a female *Anopheles* mosquito infected with one of the four *Plasmodium* species infectious for humans. The mosquito's saliva carries the malarial sporozoites into the blood. Approximately 30 minutes later these sporozoites enter the liver. Once in the liver, the sporozoites divide over the course of about 5 days, forming a schizont. A schizont may contain up to 30,000 merozoites which spill into the bloodstream the schizont ruptures. Within seconds, merozoites infect red blood cells (RBCs) and again replicate asexually, with each schizont producing up to 36 merozoites. Each time a RBC bursts and liberates progeny, other blood cells are infected. The cycle may continue until the

person dies of anemia and/or other complications. A few of the merozoites in RBCs differentiate into gametocytes, a sexual form, which, if ingested by a mosquito, are liberated from the RBCs in the mosquito stomach and subsequently mate. The progeny, sporozoites, accumulate in the saliva and the process starts again when the mosquito feeds (see 41 for review).

Malaria Prevention

Presently, there is no effective vaccine against malaria. For many years, **chloroquine** was a cheap and effective therapeutic for treating malaria, but in recent years chloroquine-resistance has increased dramatically. Indeed, a seven-fold increase in malaria in Senegal has been linked to emergence of chloroquine resistance, yet resistance in this country is much less prevalent than in other parts of Africa [44]. The same is true for newer anti-malarial drugs. Within just five years of mefloquine's arrival in Thailand, resistant parasites had emerged. In parts of Thailand and Cambodia, there are now parasites that are resistant to all the conventional drugs. Today, almost no endemic country is without drug resistant parasites. Moreover, because manufacturing such drugs tends to be unprofitable, fewer drug companies than ever before are developing new anti-malarials.

Perhaps surprisingly, one of the most effective methods for combating malaria in developing countries has been the use of **insecticide impregnated bed-nets**. Some scientists argue that such control methods – which have been shown to cut mortality and morbidity by 15 to 33 per cent in short-term trials – may in the long term only make populations more vulnerable to severe disease by reducing natural immunity. If they are correct, methods such as bed-nets and genetically engineered mosquitoes would have little impact on the control of the disease, and could indeed make matters worse [44].

Clearly, the most effective approach to combating malaria is an effective vaccine. As has been demonstrated with smallpox, a coordinated worldwide vaccination program can result in the eradication of communicable diseases

Malaria Vaccine Development

There are three recognized approaches to malaria vaccine development which are proposed to function by interrupting the parasite's lifecycle at three different stages.

The first, and most attractive approach, is the **pre-erythrocytic vaccine** which aims to block sporozoite entry into the hepatocyte and/or release of merozoites into the blood stream. Immediately following infection, sporozoites migrate to the liver and begin the exoerythrocytic stage of their lifecycle. An ability to block hepatocyte entry or the destruction of infected hepatocytes prior to liberation of merozoites would prevent the disease, the passage of the parasite on to feeding mosquitoes, and merozoite release and subsequent invasion of red blood cells.

A second approach is to develop an **'antidisease' vaccine**. This would not prevent infection, but it would block the body's harmful response to infection, which is thought to be responsible for many of the symptoms associated with infection.

A third approach, known as the **'altruistic' vaccine**, would not stop infection or symptoms in the individual but would prevent infection from spreading to others.

ICC's candidate malaria vaccine is a pre-erythrocytic vaccine - it aims to prevent infection by enabling the immune system to 'clear' the pathogen prior to the release of merozoites from hepatocytes (events up to this point are asymptomatic). A historical perspective of this type of malaria vaccine is given below.

A Brief Review of Pre-Erythrocytic Vaccine Candidates

The quest for a malaria vaccine was built on the observation that people repeatedly bitten by mosquitoes gradually acquire immunity over a period of many years [44]. Moreover, in the 1960s the British scientists Sydney Cohen and Sir Ian McGregor discovered that antibodies from such individuals could reduce the parasite load in people lacking immunity and clear the disease symptoms.

Around the same time, researchers at New York University (NYU) achieved full protection for the first time by injecting animals with small numbers of sporozoites from mosquitoes that had previously been irradiated. Later, researchers at the University of Maryland, NYU and Walter Reed Army Institute showed that 90% of a group of human volunteers immunized with sporozoites from irradiated sporozoites later resisted exposure to virulent sporozoites [6, 7]. This confirmed that protective immunity to the sporozoite stage (i.e. the pre-erythrocytic stage) of the malaria parasite could be induced [45]. However, an inability to culture sporozoites *in vitro* thwarted the possibility of using sporozoites as a vaccine. Sporozoites still cannot be cultured *in vitro*.

The strategic development of a synthetic malaria vaccine required the identification of immunodominant/neutralizing malaria epitopes. In 1985, a group at NYU led by Drs. Ruth and Victor Nussenzweig, identified the dominant B cell epitope from the circumsporozoite protein (CS), a major component of the sporozoite surface membrane at the time the parasite enters the bloodstream [5]. Antibodies to this epitope (NANP)_n were shown to be sporozoite neutralizing by protecting against rodent and human malaria [8]. Antibodies to the CS protein also appeared to correlate positively with protection in naturally infected individuals.

These studies leave little doubt as to the ability of anti-CS repeat antibodies to protect against malaria infection, providing sufficient antibody titers can be raised. The identification of this epitope therefore enabled the strategic development of synthetic CS-based malaria vaccines. Several vaccine candidates employing different carriers were developed based upon the identification of this epitope; a brief overview of four of them is given below.

1. Tetanus Toxoid Conjugated Synthetic Peptides

The (NANP)₃ synthetic peptide, conjugated to the protein carrier tetanus toxoid (TT), was the first synthetic malaria vaccine to undergo phase I and phase II clinical trials in the late 1980's [9-11]. TT is widely known to provide powerful T cell help for coupled immunogens. Of the thirty-five vaccinees, the three having the highest titers of anti-sporozoite antibody were selected for challenge studies. One of the vaccine recipients remained free of parasitaemia at 29 days, while the other two did not exhibit asexual stage parasites until 11 days, compared with a mean of 8.5 days for the un-vaccinated control group. Therefore, protection appeared to correlate positively with anti-NANP titers.

The limited effectiveness of this vaccine was attributed to suboptimal levels of anti-NANP antibodies. Attempts to increase dosage were hindered by toxicity of the TT carrier. Further, the lack of parasite-derived determinants capable of priming malaria-specific T cells also likely contributed to the low levels of protection.

2. FSV-1

Short synthetic peptides often have an *in vivo* half-life which is too short for them to be effective as prophylactic or therapeutic drugs. Standard approaches for increasing the immunogenicity of peptides is to either couple them to larger carrier proteins, or to assemble them into multimeric structures. In this case 32 copies of the CS repeat sequence ((NANP)₁₅(NVDP))₂ were linked and produced recombinantly fused to a random 32 amino acid fusion protein [12]. This vaccine candidate was called FSV-1.

Upon immunization, twelve of the fifteen volunteers developed antibodies that reacted with sporozoites. No patients exhibited adverse reactions to the protein, indicating that the NANP repeat itself is non-toxic. Of the fifteen patients immunized with 3 doses, six were selected to receive a fourth dose and were then challenged with the malaria parasite. Parasitaemia did not develop in the volunteer with the highest titer

of CS antibodies, and parasitaemia was delayed in two of the other five vaccinees. As with the NANP-TT vaccine discussed above, protection appeared to correlate positively with anti-NANP titers. This vaccine was deemed partially successful in that it reconfirmed that humans can be protected by CS protein subunit vaccines. However, the level of protection was not sufficient to warrant larger trials of this particular candidate.

The major shortfall of this vaccine was that it did not provide an efficient source of T cell help. The only individuals who would have received T cell help from this vaccine would be those in whom the CS repeat serves as both a B and Th cell epitope. However, this sequence is known to be a Th epitope for only a limited number of individuals, i.e. it is highly genetically restricted. In an attempt to circumvent the genetic restriction of using the NANP repeat alone, Good, Berzofsky and colleagues identified a superior Th epitope (326-343) on the CS protein outside the NANP repeat region, which appeared to be the immunodominant T cell site on the CS protein. When coupled to the NANP repeat, the Th epitope did indeed appear to circumvent the genetic restriction observed when NANP alone was the immunogen [13]. This epitope formed the basis for the development of the universal T cell epitope to be discussed later.

3. Multiple Antigenic Peptides

Nardin and coworkers at NYU have been able to elicit relatively high titers of anti-CS antibody in a diverse range of genetic backgrounds by combining the NANP repeat epitope with the T cell site identified by Berzofsky and Good [13] in a MAP format [14]. Using their proprietary 'universal' form of the CS-T cell epitope, Nardin and co-workers have been able to elicit anti-CS antibodies in all genetic backgrounds tested, suggesting that genetic restriction is alleviated by inclusion of this epitope.

While MAPs have proven to be excellent research tools, providing valuable insight into immune recognition of the CS protein, there are several intrinsic problems associated with using them in a commercial vaccine. Their commercial utility has yet to be established relative to manufacturing and cost issues.

4. DNA Vaccines

Over the past 5 years, significant progress has been made in the development of DNA vaccines which can potentially protect against malaria infection [42]. This class of vaccines appears to be particularly amenable to the efficient priming of CTL responses, although, as with all DNA vaccines, the exact mechanism by which this is achieved remains unresolved. In 1994, a group led by Dr. Steven Hoffman successfully protected 9 of 16 (56%) mice against *P.yoelli* infection using a DNA vaccine expressing the *P.yoelli* CS protein [15]. It was later shown that this vaccine protected only one of five strains of mice (H-2d) to a level of 75% [16]. A second DNA vaccine (PyHEP17) protected three of the five strains (H-2a, 71%; H-2k, 54%; H-2d, 26%) and the combination of vaccines protected 82% of H-2a, 90% of H-2k, and 88% of H-2d mice. Protection was absolutely dependent on CD8+ T cells, INF-gamma, or nitric oxide [16]. A human clinical trial using a similar multi-gene DNA vaccine designed to induce protective CD8+ T cell responses against *P. falciparum* infected hepatocytes was recently initiated [17, 18]. Initial results from these trials are expected sometime in 1998.

While there is only minor opposition to the use of gene therapy to treat life-threatening conditions such as cancer, its prophylactic use in healthy individuals remains controversial. These concerns are driven by the unknown long term effects of DNA immunization, such as the fear that the DNA may integrate into chromosomes to induce mutations and genetic abnormalities and the propensity for the production of anti-DNA antibodies associated with autoimmune diseases such as Lupus. There is already evidence that young children with certain predispositions could be at risk for certain forms of organ specific autoimmune problems [42]. For populations living in endemic areas, the benefits of protection against such a lethal disease as malaria may out-weigh the concerns associated with the unknown side-effects of DNA immunization. However, for travelers from North America and Western Europe, they may not.

While DNA-based therapeutics and prophylactics have clearly introduced an added dimension to modern medicine, many believe that they should not be regarded as a universal solution to all ailments. We are of this belief and remain convinced that parallel development of vaccines using alternative approaches is

essential when targeting a pathogen which continues to resist multiple vaccine strategies, almost 25 years after humans were first protected against malaria using irradiated sporozoites [6, 7].

5. HBsAg-CS

One of the most promising malaria vaccines of recent times utilizes the hepatitis B surface antigen (HBsAg) to deliver CS epitopes, an approach developed by SmithKline Beecham (SB) and colleagues. The CS epitopes include the NANP repeat, in concert with additional CS epitopes, including the T cell site identified by Berzofsky and Good [13] (but not the universal form developed by Nardin and co-workers [3, 4], fused to the hepatitis B surface protein [19]. This vaccine was recently the subject of human clinical trials [19]. When administered with one of three different adjuvants, this vaccine protected 1/7, 2/7 and 6/7 individuals respectively. Of the seven individuals immunized with vaccine 2 (adjuvant: oil-in-water emulsion), none of the five patients with anti-CS titers (IFA) in the range of 100-12,800 were protected, while the two vaccine recipients with antibody titers in the range of 25,600-51,200 were both protected. Again, protection correlated with anti-CS titers.

The strong reliance of this candidate vaccine on powerful immune-enhancing adjuvants was evident as they appear to be critical in compensating for its apparent low immunogenicity. For example, the enhanced protection to vaccine 3 (adjuvant: monophosphoryl lipid A and QS21), did not correlate well with antibody titers. This suggests that either CTL priming or non-specific immune stimulation may be playing a significant role in protection; if the latter is true protection may be short-lived. Patients were protected against malaria for 60 days, but no data has been presented on whether these patients remain protected beyond this period. This vaccine is currently undergoing field trials in the Gambia.

There are legitimate concerns as to whether such powerful adjuvants are too toxic for human use. Indeed, several participants of SB's clinical trial experienced "severe symptoms" after the second dose of vaccines 2 and 3 [19], while none of those receiving vaccine 1 (adjuvant: alum and monophosphoryl lipid A) exhibited such symptoms. These adverse systemic reactions "may have resulted from the intensity of the immune response after the second dose" [19]. However, toxicity of the adjuvant cannot be ruled out, particularly in light of the fact that no adverse reactions were noted with vaccine 1 (adjuvant: alum and monophosphoryl lipid A), and none are routinely observed with the existing hepatitis B virus vaccine which has a very similar composition. Unlike alum, the adjuvant QS21 has not been approved for human use and strong reliance on such adjuvants for vaccine efficacy may be a concern.

Table 1: Comparison of Primary Antibody Responses After Immunization with HBsAg and HBcAg.

Strain	H-2	Anti-HBs (Titer)	Anti-HBc (Titer)
B10	B	256	40,960
B10.D2	D	1024	81,920
B10.S	S	0	163,840
B10.BR	K	32	163,840
B10.M	F	0	20,480
B10.P	P	1024	10,240
C3H.Q	Q	2048	327,680
BALB/c	D	1024	327,680

Groups of mice of the indicated strains were immunized with a single dose (4µg) of either HBsAg or HBcAg and sera were examined for anti-HBs and anti-HBc antibodies 3 weeks later by ELISA) [46].

Like HBc, HBs is a particulate protein derived from the hepatitis B virus which has been proposed as a carrier for heterologous epitopes. We have studied the relative immunogenicity of HBsAg compared with HBcAg, and the ability of each to evoke immune responses in different genetic backgrounds (Table 1) [46]. These data emphasize both the higher immunogenicity of HBc relative to HBs, and the universal

responsiveness to HBc, irrespective of genetic background. For example, HBc is >300x more immunogenic than HBs in BALB/c mice; and, while both B10.S and B10.M mice are non-responders for HBs, every strain tested is responsive to HBc. These results re-emphasize the suitability of HBc as a vaccine carrier and specifically, its superiority over HBs, hence our selection of HBc as opposed to HBs to carry heterologous epitopes, and, in the case of malaria, overcome genetic restriction and inadequate antibody titers which have prevented the development of an effective vaccine.

Summary of Previous Pre-Erythrocytic Stage Malaria Vaccine

The positive correlation between protection against malaria infection and anti-CS antibody titer has been demonstrated repeatedly over the past 15 years [9-12, 19]. The evidence that a vaccine which can elicit high-titer, long-lived antibody responses in sufficient vaccine recipients can be protective suggests that protection against malaria infection is achievable via neutralizing antibody production. With delivery systems for the NANP epitope apparently exhausted, many groups have switched focus to alternative delivery systems, such as DNA and powerful adjuvants, in concert with the identification of new pathogen-neutralizing B and T cell epitopes from other stages of the parasitic lifecycle (discussed in "Phase II" section). While this work will increase the repertoire of reagents which can be used to develop effective malaria vaccines, we remain convinced that a delivery system that optimizes the interaction between the CS epitopes at hand, and the host immune system, will result in an effective malaria vaccine. The high level of protection (90-100%) achieved with the HBc/CS-repeat is highly encouraging, and clearly warrants further testing of the *P.falciparum* homolog..

The Role of Cytotoxic T cells in Pre-Erythrocytic Stage Immunity

The exact role of cytotoxic T cells in sterile immunity to malaria remains to be determined, although immunity elicited by irradiated sporozoites can be eliminated in some strains of mice by treatment with antibodies to CD8⁺ T cells. The exoerythrocytic form of the parasite found in the hepatocytes has been identified as a target of this cell-mediated immunity [Hoffman, 1989 #648]. However, HBc particles expressing just the *P.bergei* CS-repeat epitope afforded >90% protection against *P.bergei* infection [1], suggesting that priming of CD8⁺ cells may not be essential for protection. Further, the successful protection of >90% of mice against *P.yoelli* with an HBc particle expressing only the *P.yoelli* CS-repeat tends to conflict with previous suggestions that cell-mediated immunity is necessary for protection in this rodent model, providing titers of neutralizing antibody are sufficiently high [2].

The contribution of T_c cells in vaccine trials involving irradiated sporozoites, or large portions of the CS protein which includes T_c epitopes such as SB's HBsAg-CS vaccine, remains to be determined. The strength of the HBc carrier technology is in the delivery of defined, linear B cell epitopes. Although HBc has been shown to prime pathogen specific T_h cells efficiently by independent groups [1, 20], HBc has not been shown to deliver foreign CTL epitopes, although a CTL response to HBc is elicited during the natural course of HBV infection [21, 22], and is the focus of new HBV vaccines [23-25]. These studies have shown that optimal CTL effects of HBc-CTL epitopes appear to require alternate deliver mechanisms, such as DNA [25, 26] and complex adjuvant formulations [23].

We believe strongly that there is sufficient data to suggest that immunity to malaria can be achieved through neutralizing antibody specific for the NANP repeat, providing that antibody titer is sufficiently high and long-lived in the majority of vaccine recipients. While we are confident that the HBc carrier, the NANP epitope and CS universal T cell site will afford such immunity, we remain cognoscente of the benefit of incorporating B and T cell epitopes from other malaria proteins expressed during other stages of the parasitic lifecycle, into our vaccine (see Phase II work).

Hepatitis B Core Protein

The hepatitis B virus core protein (HBc) is a 22kDa protein which spontaneously assembles into a particulate structure (27 nm) in the course of virion assembly during HBV infection and also during heterologous expression in both Prokaryotic and Eukaryotic systems [27]. Unlike the surface protein of the hepatitis B virus (HBs), HBc is highly immunogenic and immune responses to it show no significant degree of genetic restriction (see Figure 1) [46], making it an ideal candidate to serve as a carrier moiety.

In the late 1980s it was determined, using synthetic peptides, that the major immunodominant B cell epitope on HBc is localized around amino acids 75-83 [47; Milich and Thornton, unpublished observation]. Since the identification of this site, several research groups have successfully cloned foreign epitopes into this immunodominant loop and successfully demonstrated that HBc is indeed a unique and highly effective carrier moiety for B cell epitopes[27]. The multimeric, particulate nature of the assembled HBV nucleocapsid confers this enhanced immunogenicity to foreign sequences. Experimental examination of a variety of B cell epitopes, either chemically linked or fused by recombinant methods to HBc particles, has yielded significant success [27].

Earlier this year, two research groups based at the NIH and the MRC in the UK, solved the structure of HBc particles using cryo-electron microscopy (see Figure 2) [28, 29]. For those familiar with the immunological properties of HBc, it was not surprising to learn that the region encompassing amino acids 75-83 does indeed form a loop at the surface of the particle. Specifically, amino acids 78-82 appear to form a loop that connects adjacent helices (Figure 2B). Preliminary x-ray diffraction data is consistent with these predictions (R.Crowther - MRC, Seminar given at the Scripps Research Institute, Oct.1997).

HBc as a Carrier of Heterologous Epitopes

The hepatitis B core protein is one of the most immunogenic proteins known. In its natural state it forms the core of the hepatitis B virus and the immune system of essentially every individual who becomes infected with the virus develops a powerful immune response to the core [30].

Recombinantly produced, hybrid hepatitis B core proteins have been shown to significantly enhance the immune response to inserted foreign sequences displayed at the surface with a conformation that mimics the structure of the sequence in the parent protein [27]. The variety of hybrids produced in this manner have proven HBc to be amazingly tolerant to the internal addition of foreign sequences at the site of the immunodominant loop, while retaining the ability to form particles that can be easily purified in a manner that is largely independent of the inserted sequence.

As a carrier moiety for B cell epitopes HBc offers other significant benefits. The particulate structure stimulates immunogenicity and allows incorporation of multiple epitope copies that are conformationally constrained. These features confer superior results relative to non-particulate carriers such as KLH, BCG, or tetanus and diphtheria toxoids [31]. Further, HBc can directly activate B cells, elicit strong T_H cell responses, and is efficiently processed and presented by antigen presenting cells (unpublished data, D.Milich personal communication, 1997).

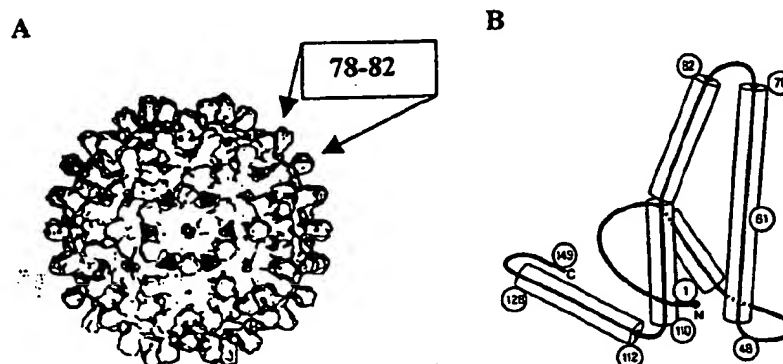


Figure 2: A: HBc particle structure determined at 9 angstrom resolution from cryo-electron micrographs. The immunodominant loop (78-82) is indicated. (From: Conway *et al.*, (1997) Nature 386, 91-94. B: Proposed polypeptide fold for HBc. Alpha-helical regions are shown as cylinders and the putative N- and C-termini are marked. An approximate numbering scheme for amino acids is given (From Bottcher *et al.*, (1997) Nature 386, 88-91)

A number of pathogen-related B cell epitopes have been chemically linked, or fused by recombinant methods to HBc, as a method to increase immunogenicity (HBV, HIV-1, FMDV, HRV-2, BLV, FeLV, HCV, MCMV, PV-1, SIV) (see 27 and 32 for review). These studies, conducted by a number of independent laboratories, have yielded significant success; including complete protection against FMDV using a HBc/FMDV particle which elicited neutralizing antibody titers comparable to those observed following exposure to the FMD virus [33]. This work proved that neutralizing epitopes, presented in the context of HBc, can elicit protective (pathogen neutralizing) immunity.

Using rodent malaria models, ICC's founding scientists successfully demonstrated that malaria CS-repeats fused to the immunodominant loop of HBc were able to protect mice against both *P.bergeii* and, perhaps more impressively, *P.yoelii* to levels of 90-100% [1, 2]. Further, antibody responses to the *P.bergeii* particle were shown to prime antibody responses effectively in a wide range of genetic backgrounds, confirming the universal priming effects of HBc (Table 2) [1].

Another advantage of the HBc carrier is the fact that it does not require complex adjuvants for efficacy. This is due to the inherent high immunogenicity of the particle. A comparison of the immunogenicity of HBc-CS1 particles showed that alum, which is approved for human use, was more effective than either IFA or CFA (Table 3) [Schodel, 1994 #682]. The importance of this observation is highlighted by toxicity problems associated with newer, more complex adjuvants as was recently noted in clinical trials of SB's candidate malaria vaccine [19].

Table 2: Immunogenicity of HBcAg-CS1 Particles in Mice.

Strain	H-2	HBcAg	CS	[DP4NPN]
C57BL	b	655,360	163,840	>655,360
B10.S	s	163,840	163,840	>655,360
B10.M	m	10,240	10,240	163,840
B10.BR	k	10,240	10,240	655,360
BALB/c	d	40,960	40,960	>655,360

Groups of three mice of the indicated strains were immunized with 20 µg of HBc-CS1 particles in CFA and boosted with 10 µg in IFA. Sera were collected before immunization, 2 weeks after secondary immunization. Sera were pooled and analyzed by solid-phase ELISA: HBcAg (50ng/well); CS represents *P.bergeii* sporozoites (1,000/well); and the *P.bergeii* repeat [DP4NPN]2 (1.0 µg/well) were the solid phase ligands. [1] (N.B. sporozoites were limiting for CS titer determination; Florian Schodel, personal communication).

Table 3: Effect of Adjuvant on Immunogenicity of HBcAg-CS1 Particles.

Immunogen	Adjuvant	Time weeks	Anti-[DP4NPN]
HBcAg-CS1	CFA	2	10,240
		4	163,840
	IFA	2	2,560
		4	163,840
	Alum	2	10,240
		4	655,360

Groups of three BALB/c mice were immunized with HBcAg-CS1 particles (10 µg) prepared in three different adjuvants; CFA, IFA or alum. Sera were collected before immunization and 2 and 4 wk after primary immunization. Sera were pooled and analyzed by solid-phase ELISA using [DP4NPN]2 (1.0 µg /well) as solid-phase ligands. [1]

The B cell Epitope, (NANP)_n

The immunodominant B cell epitope of the CS protein of *P.falciparum* is a highly conserved, repetitive tetrapeptide (NANP) [5], and antibodies to this epitope have been shown to be sporozoite neutralizing by protecting against rodent and human malaria. Immune responsiveness to this epitope has been positively correlated with immunity to malaria in both vaccine recipients and naturally infected individuals. Indeed, a review of clinical trials data for pre-erythrocytic vaccines described previously (HBs-CS, FSV-1, NANP-TT) highlights strong correlation between antibody titer and protection [9-12, 19]. Those individuals who have been protected by previous vaccine candidates are those with the highest anti-NANP antibody titers, with the possible exception of SKB's candidate vaccine (# 3) where adjuvants appeared to play a critical role in protection [19].

The Company believes that HBc is indeed the 'superior carrier' which will enable the development of high titer, long-lived antibody responses in vaccine recipients. Further, we believe that inclusion of epitopes which enable malaria specific T_H function will significantly enhance the potency and specificity of the vaccine.

In 1993 ICC founding scientists reported the protection of mice against *Plasmodium bergerii* infection using a vaccine constituting the *P.bergerii* repeat presented at the surface of HBc particles (HBcAg-CS1) [1]. More recently protection against *P.yoelli* has been described using a similar approach [2]. Initial evaluation of a particle displaying epitopes from *P.falciparum* were encouraging, but antibody titers in mice were lower than those observed for the *P.bergerii* and *P.yoelli* particles.

Table 4: Comparison of Immunogenicity of HBcAg-CS2 and HBcAg-CS2.1 in Mice. [1]

Immunogen	Strain	Time (days)	Antibody Titer (I/dilution)	
			HBc	[NANP]
HBcAg-CS2	B10	10	10,240	2,560
		24	40,960	10,240
		Secondary	655,360	655,360
	B10.S	10	10,240	640
		24	40,960	1,280
		Secondary	>655,360	20,480
	BALB/c	10	10,240	0
		24	40,960	640
		Secondary	>655,360	40,960
HBc-Ag-CS2(new)	B10 x B10.S F1	10	1,280	5,120
		24	10,240	40,960
		Secondary	163,840	2,621,440

Groups of three mice each of the indicated strains were immunized with 20 µg of hybrid HBc-CS2 or HBc-CS2.1 particles in CFA and boosted with 10 µg in IFA. Sera were collected before immunization, 10 and 24 d after primary and 2 wks after secondary immunization. Sera were pooled and analyzed by solid-phase ELISA: HBcAg (50 ng/well); the *P.falciparum* repeat [NANP]₅ (1.0 µg/well). [1]

Scientists at ICC have recently improved the immunogenicity of the *P.falciparum*-HBc particle by altering the insertion strategy of the NANP repeat epitope at the surface of the HBc particle (HBc-CS2(new)) (proprietary data). This approach has yields a more immunogenic particle and evokes antibody titers which approach those achieved for the protective *P.bergerii* particle (Table 4) [1]. We are unaware of anti-NANP titers of this magnitude being achieved following a 2 dose immunization

regime. In addition to increasing the immunogenicity of the NANP repeat, we have simultaneously reduced the HBc-immunogenicity (Table 4). A direct comparison of the two particles, using identical mouse strains, is currently underway. The poor response of CS2 in BALB/c mice is somewhat puzzling since the anti-HBc response was very high and HBc does not have a history of exhibiting genetic restriction. However, we anticipate that this can be overcome during phase I work by optimizing the presentation of the NANP repeat on the HBc particle using the new presentation strategy and the inclusion of the malaria specific universal Th epitope which is highly effective in this strain (described below) (Table 7).

The Malaria Specific T cell Epitope, CS 329-345

While the T cell help afforded by HBc is highly effective in enhancing antibody responses (i.e. B cell mediated) to 'carried' epitopes following vaccination, it will not activate malaria specific T cells. To ensure the priming of malaria specific T-helper cells as well as B cells, we will include a malaria specific T-helper epitope in our vaccine.

Table 5: Comparison of Proliferative Responses of T cell Clones after in vitro challenge with peptide 326-345 variants of the *P.falciparum* CS Protein.

Variation in Sequence 326-345	Response				
	DR9-CA2F9	DR7-RMB11	DR7-RM2B10	DR4-DW2F9	DR1-RM1B10
EYLNKIQNSLSTEWSPCSVT	+++	+++	+++	+++	+++
K K	+++	++++	+++	+++	-
K KR	+++	+++	+++	+++++	-
K KT	++++	+	+++	-	-
K KT K	+	-	+	-	-
K Q	++++	++++	++	++	-
K Q K	++	+++	+	-	-
K Q R	+	+++	-	-	-
Q K	++++	++++	++	+++	-
Q KT K	+	++	+	-	-
Q K K I	++	++	++	-	-

The data are presented as percentage of stimulation obtained with 1ug/ml control peptide NFS4 326-345. -, <20%; +, 20-49%; ++, 50-79%; +++, 80-120%; +++, 120-200%; +++++, >200%. After in vitro challenge with peptide 326-345 at 1ug/ml the stimulation index for DR9-CA2F9 was 82, for DR7-RMB11 was 30, for DR7-RM2B10 was 60, for DR4-DW2F9 was 6, and for DR1-RM1B10 was 47. [4]

The benefits of the inclusion of a universal T_H cell epitope derived from the malaria parasite are several-fold. First, the priming of malaria specific T_H cells ensures that, should a vaccine recipient be exposed to malaria, a more rapid and stronger anti-malaria response will be activated due to previous priming of malaria specific T-helper cells. Secondly, vaccinees living in malaria endemic regions will experience natural 'boosting' every time they are exposed to the parasite because their immune systems have been primed at both the B and T_H cell level. This effect is similar to clinical boosting by re-vaccination – a process which would be difficult to enforce in third-world countries where malaria is often endemic.

While the CS gene is largely invariant, limited sequence variation has been noted to occur mainly in the immunodominant T cell epitopic domains. The fact that genetic mutations always appear to result in amino acid substitutions suggests that pressure at the protein level, possibly immunological pressure, has selected for variation. Typically, the problems associated with amino acid variability of an epitope can only be resolved by the inclusion of multiple variants of the epitope. However, Nardin and coworkers at NYU recently identified a consensus form of the T cell epitope CS 326-345 which appears to bind to all class II molecules [3, 4]. Studies have shown that this consensus epitope is 'universal', like the T cell help afforded by HBc, and suggests that it will prime malaria specific Th cells in essentially all vaccine recipients (see Table 5). The fact that this epitope of the CS protein was identified by CD4+ T cells of volunteers protected against malaria following exposure to irradiated sporozoites confirms that this epitope is efficiently processed

and presented *in vivo* by antigen presenting cells (APC) when presented in the context of sporozoites [34]. The identification of this epitope was a significant advancement in the task of developing a pre-erythrocytic stage malaria vaccine, and it is this 'universal' T cell epitope that we are incorporating into our pre-erythrocytic stage malaria vaccine.

Table 6: Mapping of Epitopes within Amino Acid Sequence 326-345 recognized by T cell Clones Restricted by Different DR Alleles.

Amino Acid Sequence	Stimulation Index											
	DR9-CA2F9			DR7-RM2B10			DR4-DW2F9			DR1-RM1B10		
	10 ^a	1	0.1	10	1	0.1	10	1	0.1	10	1	0.1
EYLNKIQNSLSTEWSPCSVT	112	76	27	200	169	88	122	29	25	43	3	1
EYLNKIQNSLSTEWSPCS	148	114	62	226	165	53	114	9	1	93	1	1
EYLNKIQNSLSTEWSP	130	98	23	205	167	75	35	2	1	1	1	1
EYLNKIQNSLSTEW	134	102	42	1	1	1	94	9	2	1	1	1
EYLNKIQNSLST	5	1	1	1	1	1	4	1	1	3	1	1
LNKIQNSLSTEWSPCSVT	114	75	26	220	159	53	1	1	1	1	1	1
KIQNSLSTEWSPCSVT	50	6	3	196	109	8	1	1	1	1	1	1
QNSLSTEWSPCSVT	4	4	1	162	14	1	1	1	1	2	1	1
SLSTEWSPCSVT	1	1	1	189	190	82	1	1	1	1	1	1
STEWSPCSVT	10	4	3	7	1	1	1	1	1	1	1	1

Cloned T cells (2×10^4) were cultured with irradiated autologous PBL (5×10^4) in the presence of various peptide concentrations ranging from 0.1 to 10 μ g/ml. IL-2 levels were measured in 24-h supernatants using a bioassay based on proliferation of an IL-2 dependent T cell line.^b Peptide concentration (μ g/ml). [4]

Table 7: Genetic Restriction of Response to MAP Containing Universal T_H Epitope.

Responder	Strain	H-2	ELISA Titer
High	BALB/c	D	248,335
	DBA/1	Q	137,772
Intermediate	C57BL/10	B	57,926
	RIIIS	R	34,443
	A/J	A	23,525
Low	SJL	S	4,302
	C3H	K	3,620
	P/J	P	1,280

Results shown as geometric mean titers in sera obtained 20 days after the third injection of MAP using homologous MAP as Ag in ELISA. [4]

This epitope has also been shown to contain CTL epitopes recognized by human CD4⁺ CTL [4, 34, 35] as well as human CD8⁺ T cells [36] and, thus, may represent a universal T cell epitope, not only in terms of MHC binding but also in terms of immunologic function. Preliminary studies have shown that MAPs containing this epitope can elicit class-I restricted T cell responses in mice [4].

Production Issues

The initial failure of simple vaccine approaches to yield an effective vaccine against malaria has resulted in the development of increasingly more elaborate vaccine candidates, such as multiple antigenic peptides (MAPs), DNA vaccines, and engineered IgG fragments. While these approaches have shown promise, they do pose production complications, and in some cases safety concerns. Further, more elaborate vaccines are often so expensive that even if they are successful they are cost-prohibitive for commercial use,

particularly in the developing world where the vaccine is needed most.

In addition to the many functional advantages of HBc over other carriers, HBc particles are produced recombinantly in a highly cost-effective *E.coli* expression system. The use of a single DNA clone to produce a recombinant protein ensures absolute uniformity and reproducibility which cannot always be assured by chemical approaches which are often restricted by uncontrollable 'side-reactions'. Further, we anticipate that the immunogenicity enhancing properties of HBc will enable a 2 dose immunization regime for this vaccine.

Production of recombinant proteins in *E.coli* is a highly cost effective process. Presently we are able to produce in excess of 100mg of highly purified HBc/NANP particles (>98% by SDS-PAGE) from a 1L fermentation performed in a shaker flask. We are confident that yields can be improved significantly through the use of more efficient fermentation conditions. All purification procedures used to date at the laboratory scale are simple and scalable, permitting large quantities of material to be made and purified economically. The Company will out source all manufacturing of materials for toxicological and clinical testing to PrimaPharm in San Diego, CA. PrimaPharm possesses 20 years of cGMP manufacturing expertise and has the capability of producing, purifying, and vialing material for the malaria vaccine studies. It boasts major multinational pharmaceutical companies among its clients, including SmithKline Beecham and Allergan, and offers competitive pricing which will ensure that the final price of the vaccine will be highly competitive. An estimation of the cost for a 150L fermentation, plus purification and vialing, has been determined to be \$17,000. A chart to show the production cost for 20µg doses of recombinant HBc/NANP particles is presented in Table 8. PrimaPharm is located approximately 100 yards from ICC's research labs which will enable their production staff to work closely with ICC's research and development team during the scale-up process.

Table 8: Costing of ICC's malaria vaccine based on a 150L fermentation, purification and vialing at a cost of \$17,000. Yields of HBc-CS2(new) from a 1L culture grown in a shaker flask is typically 80-120 mg of purified particles. We are confident that yields closer to 200mg/L will be achievable with a production scale fermenter where pH, temperature, aeration and nutrient supply are more effectively controlled. The hepatitis B vaccine (HBsAg) is administered at doses of 10-20µg.

Yield (mg/L)	Yield from 150L	20µg Doses	Cost Per 20µg Dose
25	3,750 mg	187,500	9.1c
50	7,500 mg	375,000	4.5c
100	15,000 mg	750,000	2.3c
200	30,000 mg	1,500,000	1.2c

While toxicological studies of HBc-based products have not been conducted to date there is no evidence of toxicity due to HBc in HBV asymptomatic chronic carriers or in HBcAg-expressing transgenic mice.

Relationship with Future Research and Development

In addition to key *P.falciparum* epitopes, ICC is also in the process of licensing neutralizing CS-epitopes specific to other strains of human malaria. *P.vivax* is the second most important malaria target, and we intend to focus on the development of a *P.vivax* vaccine starting in 1999. The multimeric nature of the HBc particle means that it will be possible to assemble a multivalent malaria vaccine displaying neutralizing epitopes from both *P.vivax* and *P.falciparum*, either by co-expressing different HBc genes [37], or by inserting multiple epitopes into a single HBc gene. Multivalent vaccines are highly desirable as they enable costs, from production through vaccine administration, to be reduced significantly.

ICC is also actively involved in identifying additional human and veterinary vaccine targets using identified, neutralizing, linear epitopes which are amenable with presentation by HBc. We expect to identify our next 2 targets by the 2nd quarter of 1998 and begin pre-clinical development shortly thereafter. Our

experiences with the development of a HBc-malaria vaccine has already afforded us with additional information on to select candidate epitopes and expedite the development of HBc hybrid particles at the research level. We anticipate that this commonality in products will be even more important once we advance to the production stage and on to clinical testing.

Phase II

The phase II section of this proposal will focus on the clinical development of the vaccine once the lead has been identified at the end of phase I. Our clinical procedures will follow the "Guidelines for the Evaluation of Plasmodium Falciparum Vaccines in Populations Exposed to Natural Infection" distributed by the World Health Organization. The time-line for clinical development of the malaria vaccine is outlined below.

4 th Quarter 1998	Identify Lead Compound (Phase I)
4 th Quarter 1998	Identify Clinical Sites
1 st Quarter 1999	Develop Manufacturing and Purification Methods for cGMP Material
2 nd Quarter 1999	Complete GLP/GMP Manufacturing of Vaccine
3 rd Quarter 1999	Complete Toxicological Studies
4 th Quarter 1999	Submit IND
1 st Quarter 2000	Phase I/II Clinical Trials

At a research level, work on the development of a next generation malaria vaccine will continue during phase II by improving the immunogenicity of the *P.falciparum* particle via the inclusion of addition epitopes. Indeed, ICC is in the process of establishing collaborative arrangements with academic groups to focus on the identification of such epitopes, including those derived from proteins expressed during liver and blood stages of the parasite's lifecycle. Potential candidates include the merozoite surface protein-1 (MSP-1), which is considered a 'blood stage antigen'. Epitopes within MSP-1 are the target of antibodies which inhibit erythrocyte invasion *in vitro* [Blackman, 1990 #484; Chang, 1992 #258; Cooper, 1992 #2309; Hui, 1994 #716], and are particularly promising candidates because they show no amino acid variation. Vaccination experiments with the equivalent polypeptide from *P.yoelii* is protective [38, 39], and protection appears to be mediated largely by antibody [38, 40], making it an ideal candidate for presentation by HBc once neutralizing epitopes have been more precisely mapped. At the present time, these epitopes are poorly defined beyond the clear importance of the highly disulfide bonded C-terminal 19kDa region of MSP-1 [Ohta, 1997 #2134; Egan, 1997 #2405; Guevara Patiño, 1997 #2408]. Once new, pathogen neutralizing epitopes derived from MSP-1, or other target proteins which are amenable with the HBc presentation strategy have been identified, they will be tested.

Key Patents Held By ICC

ICC has exclusive or semi-exclusive licenses to the following patents which are relevant to this work:

1. T Cell Epitopes for the Hepatitis B Virus Nucleocapsid Protein (Patent # 5, 143,726)
Inventors: Thornton, Moriarty, Milich, and McLachlan.
Immune Complex Corporation holds an exclusive license of this patent from The Scripps Research Institute, La Jolla, CA.
2. Immune Complex Corporation is in the final stages of negotiating exclusive and non-exclusive licenses to issued patents and patent applications describing critical malaria B and T cell epitopes necessary for the development of this vaccine. The final terms are scheduled to be completed by January 31, 1998. At the time of review, confirmation of this arrangement will be available upon request.

Potential For Commercial Application

With 7 million U.S. citizens traveling to malaria endemic areas, the initial retail market for the vaccine in the U.S. alone is over \$1.1 billion dollars annually (CDC, Harvard School of Public Health). The market size in dollars for purchases by point of care for a 50% market share will be closer to \$250 million annually.

In addition to the US travelers market, the market in dollars of travelers from other developed countries to malaria endemic areas is thought to be at least twice that of the U.S. market or another \$500 million. A total of 30-35 million people from developed countries travel to less developed ones, many of which are in the tropics. At least 30 % of those succumb to illness [47]. As many as one in every 100 people from industrialized countries who visit West African countries will get malaria; those who visit rural areas, away from the relative safety of air-conditioned urban hotels, are the most likely to become infected.

Sales to the World Health Organization at a largely reduced price but high volume is estimated at \$20 million annually (60c per dose; Howard Engers, personal communication with Ben Thornton, 1997). Malaria is endemic in 91 countries. Eighty percent of the cases occur in Africa where it accounts for 10% to 30% of all hospital admissions. Malaria, thus, is a heavy economic burden on developing countries. The cost of treatment is between \$US 0.08 and \$US 5.30 depending on the drugs prescribed. The total cost of malaria in terms of health care, treatment, lost production, etc. is estimated to be \$US 1.8 billion. For example, India will spend U.S. \$40 million on malaria control this year – up 60 % from last year. It is also planning a five-year, \$200 million program targeting 210 million people in 100 high-risk districts. The program will introduce new tools such as pesticide-impregnated bednets – a much less-effective preventative measure compared with an effective vaccine [44].

Although other vaccines are currently being developed by other pharmaceutical companies, the Company believes that its HBc-based malaria vaccine will have economic, compliance, and efficacy advantages over its competition.

B. RELEVANT EXPERIENCE: PI, STAFF, and CONSULTANTS

Ashley J. Birkett, Ph.D. (Principal Investigator)

Dr. Birkett has extensive experience working with the hepatitis B core protein. He is particularly familiar with the use of hepatitis B virus core protein (HBc) as a carrier of foreign peptide sequences. These include the insertion of foreign epitopes for raising polyclonal antibody, expression of short proteins as C-terminal extensions of HBc, and antigenic/immunologic characterization of hybrid HBc particles. Dr. Birkett is the recipient of a phase I SBIR grant to study the chemical insertion of synthetic epitopes into HBc using native chemical ligation of protein fragments (Title: Epitope Scanning Using Protein Semisynthesis, Grant # 1-R43-GM54461-01), and is in the process of filing multiple patents to cover the novel uses and significant improvements he has made to the HBc carrier system. He has also worked extensively on a NIEHS contract (Contract # NIH-ES-52002, PI Dr. H.J.Barnes), currently in phase II, to use HBc as a carrier for generating high affinity, inhibitory antibodies to human cytochrome P450s for use in *in vitro* testing of candidate therapeutic drugs by drug discovery groups.

Ph.D. Biochemistry (1994)

Medical College of Virginia/Virginia Commonwealth University
Richmond, Virginia.

B.Sc. (Honors) Applied Biological Sciences (1990)

Bristol Polytechnic, Bristol, UK.

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George B. Thornton, Ph.D. (President/CEO)

Dr. Thornton joined the ICC/Synthetic Genetics in 1996, becoming the President and CEO in August. He was formerly the Vice President of Research and Development at GeneMedicine (Houston), a gene therapy biotechnology company developing non-viral approaches to gene delivery. While at GeneMedicine he was responsible for all research activities and the development (preclinical and clinical) of their gene therapy products. Two GeneMedicine programs directed by Dr. Thornton have received phase I approvals this year. Prior to his position at GeneMedicine, Dr. Thornton was Director of Research at the Johnson & Johnson PRI laboratories in La Jolla where he was responsible for research and operations at that site. While at Johnson & Johnson, scientists lead by Dr. Thornton were the first to successfully obtain a human antibody derived from phage displaying human antibody combinatorial libraries. In addition, he set up and was responsible for the J&J laboratories in Sydney, Australia which was developing gene therapy products based on ribozyme technologies licensed from the CSIRO (Canberra). The hepatitis B core technology was developed as a collaboration between Dr. Thornton's laboratory at J&J and Dr. Milich's laboratory at The Scripps Research Institute. Dr. Thornton is a Founder of the Company and co-inventor on the Company's key hepatitis B core patents.

D. EXPERIMENTAL DESIGN AND METHODS

Cloning of Universal Malaria T_H Epitope onto HBc

In the hepatitis B virus, the HBc protein is 183 amino acids in length. Carboxyl-terminal truncations of HBc to 144 are amenable with particle assembly, and are generally preferable since amino acids 150-183 encodes an arginine-rich domain which binds viral nucleic acid. When expressed in *E.coli* this domain non-specifically binds *E.coli* mRNA. The 20 amino acid universal T_H epitope will be inserted after amino acid 149 of HBc using PAGE purified complementary synthetic oligonucleotides, designed to have an *EcoRI* single strand overhangs at the 5' end, and a *SacI* overhangs at the 3' end precluding prior restriction digestion of the insert. However, the synthetic fragment will NOT restore either restriction site as it is necessary to destroy the restriction sites for later cloning steps. Once annealed, this fragment will be directionally ligated into the plasmid pKK-HBc149-CF which has *EcoRI* and *SacI* restriction sites positioned directly after amino acid 149 of HBc. Clones will be identified by restriction endonuclease mapping, and then confirmed by DNA sequencing.

Generation of HBc/NANP Hybrid Particle Expression Vectors

To construct the hybrid HBc-(NANP)_n expression vectors complete with the universal malaria T_H epitope at the C-terminus, it will be necessary to adopt a 2-step cloning strategy. First, PAGE purified synthetic complimentary oligonucleotide fragments encoding for (NANP)₃, (NANP)₆, (NANP)₉, and (NANP)₁₂ repeat epitopes will be annealed and directionally ligated into the cloning/expression vector pKK322-HBcLI. This vector has *EcoRI* and *SacI* restriction sites introduced between amino acids 77 and 78 which is the middle of the immunodominant HBc loop. Again, pairs of oligonucleotides will be designed to have *EcoRI* single strand overhangs at the 5' end, and a *SacI* overhangs at the 3' end precluding prior restriction digestion of the insert. However, the restriction sites will be restored after cloning of the insert. The second step will be to incorporate the universal T_H epitope exchanging the C-terminal fragments of HBc

for the fragment with a fragment from which also contains the universal Th epitope. All constructs will be verified by restriction analysis and sequencing prior to expression.

Expression and Purification of HBc/NANP Hybrid Particles.

Once constructed, expression will be transformed into *E.coli*, strain TB1. Cells will be grown for 36 hour in TYN media supplemented with 1 g/L glucose and harvested by centrifugation. To purify the hybrid particles, cells from a 1L culture will be lysed in PBS by a single passage through a French pressure cell. Following clarification of the lysate, the particles will be precipitated by ammonium sulfate, resuspended in 20 mM sodium phosphate, pH 6.8, and dialyzed against the same buffer. The dialyzed material will be clarified by a brief centrifugation and the supernatant subjected to gel filtration chromatography on a Sepharose CL-4B column (see Figure 3). Particle containing fractions will be identified immunologically, subjected to hydroxyapatite chromatography, and reprecipitated by ammonium sulfate prior to resuspension, dialysis, and sterile filtration prior to storage at -70°C . Typical yields of HBc-NANP particles, from a 1L shaker flask culture, are in the range of 80-120 mg of purified particles.

Physical Characterization of Hybrid Particles

HBc hybrid particles have a characteristic elution position when analyzed by Sepharose CL-4B chromatography (see Figure 3). In addition to Sepharose CL-4B chromatography, the production of uniform hybrid particles will also be confirmed by electron microscopy (see inset on Figure 3).



Figure 3: Elution profile of Hybrid HBc Particles following Sepharose CL-4B Chromatography. Arrow indicates elution range of HBc particles. Inset: Electron Micrograph of Hybrid HBc Particles.

Immunization of Mice with Hybrid Particles

Mice will be immunized with 20 μg of hybrid particles in alum, and boosted 28 days later with 10 μg of hybrid particles in alum. Mice will be bled and antibody titers measured 2 and 4 weeks post-primary immunization, and 3 weeks post boost. To determine if there is MHC-specific genetic restriction, the following mouse strains will be immunized B10(b), B10.S(s), B10.P(p), B10.D2(d), and B10.BR(k).

ELISA Testing of Anti-NANP Antibodies

Antisera will be evaluated using (NANP)5 peptide to capture anti-(NANP)_n antibodies, in an ELISA format. Briefly, (NANP)5 peptide will be coated on ELISA plates, blocked, and incubated with serial dilutions of antisera. An anti-mouse/HRP conjugate, followed by TM Blue substrate will be used to detect bound antibodies.

IFA Testing of Antisera

To test the ability of antisera to bind native sporozoites, immunofluorescence assays (IFA) will be performed on pre-dried slides. Sporozoites (1000/well) will also be coated to ELISA plates for direct antibody binding assays.

Testing Functionality of Universal Malaria T_h Epitope

Although enhanced immunogenicity of HBc-NANP particles containing the malaria universal T_h cell site as compared to particles lacking it will be one indication of "helper" function, direct assays of malaria – T_h cell function will also be performed. Mice will be primed with HBc-(NANP)₃-UTC, and the ability of a

UTC peptide to recall T cell proliferation *in vitro* will be determined. Secondly, mice will be primed with the UTC peptide and "challenged" with sub-immunogenic amounts of HBc-(NANP)₃-UTC particles *in vivo*. If the UTC-primed T cells are capable of providing T cell help for NANP-specific B cells, enhanced anti-NANP antibody production should occur. This will be determined using the ELISA described above.

Stability Studies

Sterile filtered HBc particles will be incubated at 37 °C. Aliquots will be taken at 0, 1, 3, 7, 14, and 28 days, and analyzed by reducing SDS-PAGE and ELISA. The structure of the 28 day particles will be further analyzed using electron microscopy. Unless there is a clear indication of instability, the most immunogenic of the four particles tested (as determined using non-incubated particles) will be used to immunize mice following a 28 day incubation at 37°C. In the unlikely event that the most immunogenic particle is less stable, or affords a poor yield of purified particles, an alternative particle will be selected for this experiment.

E. HUMAN SUBJECTS

No human subjects or human derived samples will be utilized during phase I.

F. VERTEBRATE ANIMALS

No vertebrate animals will be used at ICC's facility. All animal work will be contracted out to HTI-BioProducts, Ramona, CA, which has NIH approved facilities.

G. CONSULTANTS

Founding Scientists: The founding scientist of ICC include David Milich (Scripps Research Institute), Florian Schodel (Merck), Ben Thornton (ICC), Ashley J. Birkett (ICC), and Darrell Peterson (Virginia Commonwealth University). With the exception of Dr. Schodel, due to his employment at Merck, all ICC founders remain actively involved in the development of the patented HBc platform.

Scientific Advisory Board: Further, ICC has recruited a highly regarded team of scientists for its scientific advisory board (Lindsey Whitton, Ph.D., Scripps Research Institute), James Bittle, Ph.D. (Retired, former President of Pitman-Moore), Fred Brown, Ph.D. (USDA, Plum Island), John Gerin, Ph.D. (Georgetown University), Ruth Nussenzweig, Ph.D. (NYU).

Process Development & Manufacturing: Prima Pharm, San Diego CA.

Preclinical Development: The Company has engaged International Pharmaceutical Services, Inc. ("IPS") (Bridgewater, NJ) to oversee the preclinical development of R15K. IPS is a consulting organization for the pharmaceutical and medical device industries worldwide. IPS will be providing expertise to the company in the areas of preclinical development, product development, quality assurance and international regulatory affairs. The principals, Drs. Paul Wray and Jim Davis, have over thirty years experience in pharmaceutical development at Lederle, Wyeth Ayerst, Ortho Pharmaceutical Corporation (Johnson & Johnson), Ortho Biotech and Schering. Additionally, Dr. Wray is an elected Fellow of the Academy of Pharmaceutical Science, where he chairs the Industrial Pharmaceutical Technology Section, and the American Association of Pharmaceutical Scientists.

Clinical Development: The Company has engaged Advanced Biologics, LLC (Lambertville, NJ) to oversee the clinical development of its products. Advanced Biologics is a full service CRO and offers clinical trial and medical management services to the pharmaceutical industry. The principal of Advanced Biologics is Dr. Michael Corrado. Dr. Corrado was formerly Senior Director of Infectious Diseases and later Vice President of Regulatory Affairs for R.W. Johnson Pharmaceutical Research Institute (a Johnson & Johnson company). He currently sits on the ICCAC planning committee as well as the Infectious Disease Society of America (IDSA) FDA Anti-infective Guidelines Committee. Dr. Corrado has extensive experience in designing and implementing clinical strategies, from the preclinical stage through NDA filings. Advanced Biologics currently provides clinical and regulatory support for a number of small

biotechnology companies as well as large pharmaceutical companies (J&J, Merck, American Home Products, and others).

H. CONTRACTUAL ARRANGEMENTS: None

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Checklist**TYPE OF APPLICATION** (Check appropriate box(es).)☒ **NEW** application. (This application is being submitted to the Public Health Service for the first time.)☐ **REVISION** of previously-submitted application number _____
(This application replaces a prior unfunded version of a new application.)☐ **CHANGE** of Principal Investigator (if applicable)
Name of former Principal Investigator _____**1. ASSURANCES/CERTIFICATIONS**

The assurances/certifications set forth below are made and verified by the signature of the OFFICIAL SIGNING FOR APPLICANT ORGANIZATION (small business concern) on the FACE PAGE of the application. Descriptions of individual assurances/certifications are found in application instructions under "Checklist." If unable to certify compliance with any item, provide an explanation and place it after this page.

• Human Subjects; • Vertebrate Animals; • Debarment and Suspension; • Drug-Free Workplace; • Delinquent Federal Debt; • Research Misconduct; • Civil Rights (Form HHS 690); • Handicapped Individuals (Form HHS 690); • Age Discrimination (Form HHS 690).

2. PROGRAM INCOME (See discussion in application instructions under "Checklist.")

All applications must indicate (Yes or No) whether program income is anticipated during the period for which grant support is requested.

☒ **No** ☐ **Yes** (If "Yes," use the format below to reflect the amount and source(s) of anticipated program income.)

Budget Period	Anticipated Amount	Source(s)

3. INDIRECT COSTS (See discussion in application instructions under "Checklist.")

Insert the rate, if known. If the applicant organization does not have a currently negotiated rate with the Department of Health and Human Services (DHHS) or another Federal agency, it must estimate the amount of indirect costs allocable (applicable) to the proposed Phase I project. That amount should be inserted in the space provided below. The

applicant organization should also be prepared to furnish financial documentation to support the estimated amount, if requested by the Public Health Service. An applicant organization may elect to waive indirect costs if it so desires.

☐ DHHS agreement, dated: _____ % salary and wages or _____ % Total Direct Costs.

☐ No DHHS agreement, but rate established with _____, dated: _____

☒ Rate negotiation pending with the National Institutes of Health.

☐ Indirect costs allocable (applicable) to this Phase I project are estimated to be \$ _____

☐ No indirect costs requested.

4. SMOKE-FREE WORKPLACE

Does your organization currently provide a smoke-free workplace and/or promote the non-use of tobacco products or have plans to do so?

☒ **Yes** ☐ **No** (The response to this question has no impact on the review or funding of this application.)

CONSULTING AGREEMENT

This Consulting Agreement (this "Agreement") is made and effective as of January 1, 1998 between Immune Complex Corporation, a California corporation ("Company"), and David Milich, Ph.D. ("Consultant").

NOW, THEREFORE, in consideration of the above facts and the mutual promises set forth in this Agreement, the parties agree as follows:

1. Services. Company agrees to engage Consultant as an independent contractor to perform the services set forth in Exhibit A attached hereto and incorporated herein (collectively, the "Services"), and Consultant agrees to render the Services under the terms and conditions set forth in this Agreement. Consultant agrees to complete the Services in a satisfactory manner. This Agreement does not limit Consultant's ability to enter into agreements to perform services with other parties, provided such agreements or services are not in conflict with, and do not interfere with, Consultant's services hereunder.
2. Term. The term of this Agreement (the "Term") shall commence on January 1, 1998 continue for 12 months until the services are completed. Either party may terminate this Agreement when either party provides the other party with at least fourteen (14) days' advance written notice of the termination of this Agreement, which notice shall specify the date of termination. In any event this Agreement shall also terminate (a) upon Consultant's death or disability; or (b) shall for any other reason cease to be qualified, permitted or licensed to perform the services required of him under this Agreement, or under any laws or the rules of any regulating body having jurisdiction over said Services.
3. Compensation. As payment for the Services, Consultant shall be entitled to compensation as set forth in Exhibit B attached hereto and incorporated herein, which shall constitute complete compensation for the Services. Except as set forth on any schedule hereto, Consultant shall not be entitled to any other compensation or benefits for the Services. Company shall not reimburse Consultant for any expenses.
4. Relationship. The parties expressly intend, agree and understand that the relationship between them created by this Agreement is that of owner and independent contractor, and does not constitute a hiring by either party. Consultant is not an employee, partner or joint venturer of Company. Consultant shall not be treated as an employee of Company for federal or state tax purposes.
5. Taxes and Withholding. The Company will not withhold from the consulting fees any amount for taxes, social security or other payroll deductions. The Company will issue to Consultant a Form 1099 with respect to Consultant's compensation paid hereunder, will report such compensation using the same form, and will comply with other applicable tax reporting requirements. Consultant acknowledges that he will be entirely responsible for payment of any such taxes, and he hereby indemnifies and holds harmless the Company from any liability for any taxes, penalties or interest which may be assessed by any taxing authority. Consultant agrees that, upon Company's request, he shall execute any and all documents necessary to reflect or evidence that Consultant has paid all taxes on the compensation paid under this Agreement.
6. Confidential Information. Consultant recognizes that Consultant's performance of services

hereunder will involve contact with information of substantial value to the Company, which is not old and generally not known in the industry, and which gives the Company an advantage over its competitors who do not know or use it, including but not limited to, techniques, designs, drawings, methods, processes, inventions, information related to or included in patents or patent applications, research and development, equipment, prototypes, sales and customer information, customer proprietary information and prospect lists, and business and financial information relating to the business, products, practices and techniques of the Company and certain non-public information provided to the Company by its customers or business partners (collectively, "Confidential Information"). Consultant will at all times regard and preserve as confidential such Confidential Information obtained by Consultant from whatever source and will not, either during Consultant's service to the Company or thereafter, publish or disclose any part of such Confidential Information in any manner at any time, or use the same except on behalf of the Company, without the prior written consent of the Company. Further, both during Consultant's service hereunder and thereafter, Consultant will refrain from any acts or omissions that would reduce the value of such Confidential Information to the Company.

7. Disclosure of Inventions. Consultant agrees to promptly disclose in writing to the officials designated by the Company to receive such disclosures, complete information concerning each and every invention, discovery, improvement, device, design, apparatus, practice, process, method or product (collectively, "Inventions"), whether Consultant considers them patentable or not, made, developed, perfected, devised, conceived or first reduced to practice by Consultant, either solely or in collaboration with others, during the period of Consultant's service to the Company, and up to and including a period of one (1) year after termination of Consultant's services hereunder, whether or not during regular working hours, relating either directly or indirectly to the business, products, practices or techniques of the Company, or to the Company's actual or demonstrably anticipated research or development, or resulting from any work performed by Consultant for the Company.

8. Non-Solicitation. In order to protect the Confidential Information of the Company and avoid injury to the Company, Consultant agrees that during the term of this Agreement and for one (1) year thereafter, he will not, either directly or through others solicit or attempt to solicit any employee, consultant or independent contractor of the Company to terminate his or her relationship with the Company in order to become an employee, consultant or independent contractor to or for any other person or entity.

9. Assignment of Inventions. Attached hereto as Exhibit C is the Scripps Research Institute Uniform Consulting Agreement Provisions (the "Scripps Uniform Provisions"). Subject to the Scripps Uniform Provisions, Consultant hereby agrees that any and all Inventions made, developed, perfected, devised, conceived or reduced to practice by Consultant during the period of Consultant's service to the Company, and any other Inventions made, developed, perfected, devised, conceived or reduced to practice by Consultant during said period of one (1) year after termination of Consultant's services hereunder, relating either directly or indirectly to the business, products, practices or techniques of the Company or the Company's actual or demonstrably anticipated research or development, or resulting from any work performed by Consultant for the Company, are the sole property of the Company, and Consultant hereby assigns and agrees to assign to the Company, its successors and assigns, any and all of Consultant's rights, title and interest in and to any and all such Inventions, and any patent applications or Letters Patent thereon.

10. Right to Use or Publish. Nothing in this Agreement shall limit or be construed to limit Consultant's right to use or publish information which: (a) was in the public domain before Consultant's service hereunder commenced; (b) was known to Consultant to be free from any claim of other third parties before Consultant's service hereunder; (c) was developed or acquired independently of the Company; or (d) becomes public knowledge without breach by Consultant of any obligations of confidence to the Company.

11. Further Cooperation. Consultant will, at any time during Consultant's service hereunder or thereafter, upon request and, at \$ 100.00 per hour, do all lawful acts, including the execution of papers and oaths and the giving of testimony, that in the opinion of the Company, its successors and assigns, may be necessary or desirable for obtaining, sustaining, reissuing or enforcing Letters Patent in the United States and throughout the world for any and all of said Inventions, and for perfecting, recording and maintaining the title of the Company, its successors and assigns, to the Inventions and to any patent applications made and any Letters Patent granted for the Inventions in the United States and throughout the world.

12. Keeping of Records. Consultant will keep complete, accurate and authentic accounts, notes, data and records of any and all of the Inventions in the manner and form requested by the Company. Such accounts, notes, data and records, including all copies thereof, shall be the property of the Company, and, upon its request, Consultant agrees to promptly surrender same to the Company, or if not previously surrendered, Consultant will promptly surrender same to the Company at the conclusion of Consultant's services hereunder to the Company.

13. Surrender of Materials. Within three (3) days of the termination of this Agreement, or at Company's request, Consultant agrees to return to the Company all Company documents (and all copies thereof) and other Company property in his possession, or his control, including, but not limited to, Company files, notes, drawings, records, business plans and forecasts, financial information, specifications, equipment, notes, documents, memoranda, reports, files, books, correspondence, lists or other written or graphic records and the like relating to Company's business which are or have been in Consultant's possession and control, and any other materials of any kind which contain or embody any proprietary or Confidential Information of the Company (and all reproductions thereof), whether or not created by me, or which come into Consultant's possession by reason of Consultant's service to the Company, and Consultant agrees further that all of the foregoing are the property of the Company.

14. Prohibition of Misappropriation from Others. Consultant agrees that Consultant will not disclose to the Company, use, or induce the Company to use, any invention or confidential information belonging to any third party.

15. Imposed Obligations. Consultant understands that the Company may enter into agreements or arrangements that may be subject to laws and regulations which impose obligations, restrictions and limitations on it with respect to inventions and patents which may be acquired by it or which may be conceived or developed by employees, consultants or other agents rendering services to it. Consultant agrees that Consultant shall be bound by all such obligations, restrictions and limitations applicable to any Invention conceived or developed by Consultant during the period of Consultant's service to the Company, and Consultant shall take any and all further action which may be required to discharge such obligations and to comply with such restrictions and limitations.

16. No Inconsistent Agreements. Consultant represents and affirms that Consultant has no agreement with any other party that would preclude Consultant's compliance with Consultant's obligations under this Agreement.

17. Successors. The provisions of this Agreement shall inure to the benefit of, and be binding upon, the parties' heirs, personal representatives, successors and assigns. However, Consultant may not assign or delegate Consultant's obligations under this Agreement.

18. Equitable Relief. Consultant understands and agrees that, because of the unique nature of the Confidential Information, the Company will suffer irreparable harm if Consultant fails to comply with any of Consultant's obligations under this Agreement, and monetary damages will be inadequate to compensate the Company for such breach. Accordingly, Consultant agrees that the Company shall, in addition to any other remedies available to it at law or in equity, be entitled to injunctive relief to enforce the terms of this Agreement, without the necessity of posting a bond or undertaking.

19. Governing Law. This Agreement is made in San Diego, California and shall be construed and interpreted in accordance with the internal laws of the State of California. Any controversy or claim arising out of or relating to this Agreement or the breach thereof, whether involving remedies at law or in equity, shall be adjudicated in San Diego, California.

20. Attorneys' Fees. In any controversy or claim arising out of or relating to this Agreement or the breach thereof, which results in a legal action, proceeding or arbitration, the prevailing party in such action, as determined by the court or arbitrator, shall be entitled to recover reasonable attorneys' fees and costs incurred in such action.

21. Entire Agreement. This Agreement constitutes the entire agreement between the parties, it supersedes any prior or contemporaneous agreement concerning assignment of intellectual property, patent rights or trade secrets, and may be waived, modified or amended only by an agreement in writing signed by the undersigned and the President of the Company.

22. Severability. The invalidity or unenforceability of any particular provision of this Agreement shall not affect the other provisions hereof, and this Agreement shall be construed in all respects as if such invalid or unenforceable provision were omitted.

23. Miscellaneous. No covenant, term or condition of this Agreement or breach thereof shall be deemed waived unless the waiver is in writing, signed by the party against whom enforcement is sought, and any waiver shall not be deemed to be a waiver of any preceding or succeeding breach of the same or any other covenant, term or condition. The normal rule of construction, to the effect that any ambiguities are to be resolved against the drafting party, shall not be employed in the interpretation of this Agreement. This Agreement may be executed in counterparts, each of which shall be deemed an original, but all of which together shall constitute one and the same instrument. Headings or captions of paragraphs or sections of this Agreement are for convenience of reference only and shall not be considered in the interpretation of this Agreement.

24. Arbitration. Unless otherwise provided herein, any controversy or claim arising out of or relating to this Agreement shall be submitted to arbitration in accordance with the Commercial Rules of the American Arbitration Association, and judgement upon the award rendered by the arbitrator may

be entered in any court having jurisdiction thereof. Each party shall select an arbitrator, at its sole cost and expense, and such two arbitrators shall together select a third arbitrator, which third arbitrator shall be the sole arbitrator of any controversy or claim arising hereunder. The cost of expense of such third arbitrator shall be borne equally by the parties hereto. Notice of intent to submit a dispute to arbitration shall be given to the other parties as prescribed in Section 27 below.

25. Notices. All notices and other communications required or permitted to be given under this Agreement shall be in writing and shall be deemed to have been given if delivered personally or telecopied (with receipt of delivery) or three (3) days after being sent by certified mail, return receipt requested, postage prepaid, to the parties at the following addresses or to such other address as either party to this Agreement shall specify by notice to the other:

If to Company: Immune Complex Corporation
3347 Industrial Court
San Diego, California 92121
George B. Thornton, President

If to Consultant: David Milich, Ph.D.

IN WITNESS WHEREOF, the parties have caused this Agreement to be executed effective as of the date set forth above.

COMPANY

IMMUNE COMPLEX CORPORATION

By: 

George B. Thornton, President

CONSULTANT

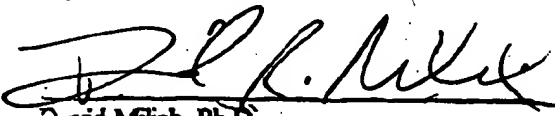

David Milich, Ph.D.

Exhibit A

SERVICES

Consultation for the Company on the hepatitis B core antigen as an epitope carrier.

Exhibit B

COMPENSATION

Consultant shall be paid at the rate of \$5,000.00 Dollars per year for services rendered under this agreement.

**IMMUNE
COMPLEX
CORPORATION**

March 23, 2000

George B. Thornton, Ph.D.
Chief Executive Officer
Immune Complex Corporation
3347 Industrial Court
San Diego, California 92121

Dear Ben,

Please provide me with copies of the two patent applications regarding (1) the stabilization of HBcAg-hybrid particles, and (2) the production of the malaria vaccine (i.e. PF 3.1). As you are aware, I have asked you and Ashley previously about the status of these applications and when I would be able to provide my input.

I designed all the antigenicity, serology and immunology experiments, provided and analyzed all the antigenicity, serology and immunology data, which were integral and necessary to the discoveries. The immunologic analysis allowed Ashley and I to constantly design, test and improve one construct after another, which ultimately led to the production of stable HBcAg-hybrid particles and to the design of the final vaccine candidate (i.e. PF 3.1). I should at least proof read these two applications before they are submitted to the patent office. I assume this is routine for an inventor on a patent. I am sure the fact that I have not been shown these applications to date is a mere oversight. My concern is the past history regarding the matter of the two *P. falciparum* SBIR grant applications. You have acknowledged on multiple occasions that I should have been listed as a co-investigator on these applications and I had requested that I be listed as a co-investigator on these applications, yet the company failed to do so.

Because I will need time to review the applications and I realize the applications are about to be submitted, I would appreciate receiving this material as soon as possible (i.e. within the next few days).

Sincerely,



David R. Milich, Ph.D.
CSO, ICC

Cc: Ed Gamson, Patent Attorney
Ashley Birkett, Ph.D., Director of Research, ICC

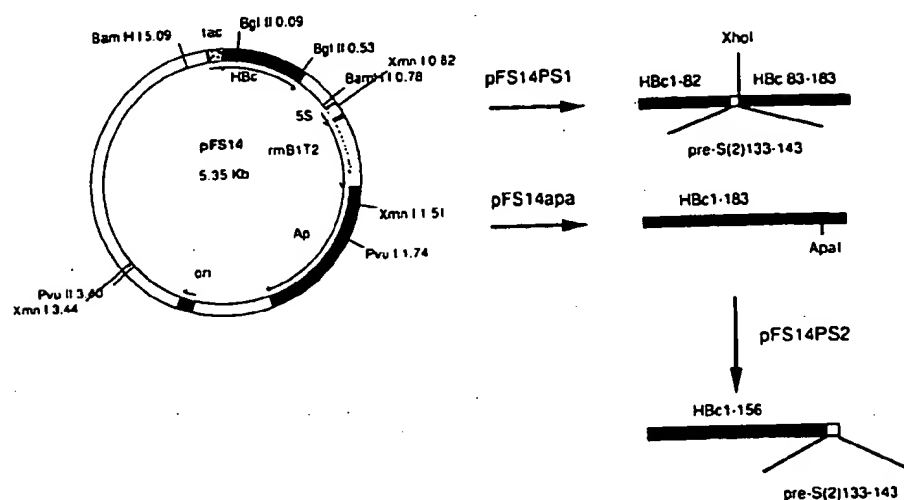


Figure 1
Structure of expression plasmid pFS14 and derivatives pFS14PS1, pFS14apa, and pFS14PS2. For details, see text.

restriction sites were bridged with a pair of complementary asymmetric synthetic oligonucleotides of the *ayw* sequence. The sequence of the coding oligonucleotide was 5'-AAT TAT GGA CAT-3'. The construct places the core gene ATG at position +11 relative to the Shine-Dalgarno sequence of pKK223 (pFS14, Fig. 1) and contains the complete nucleotide sequence of the *ayw* HBc gene. When induced with IPTG in *E. coli* JM105, this plasmid directs the synthesis of the 21-kD HBcAg immunoreactive with anti-HBcAg antibodies in immunoblots (see Fig. 3b in Schödel and Will 1989) and in ELISA of up to approximately 1% of whole-cell proteins, as judged from Coomassie-stained polyacrylamide gels (not shown). The particulate nature of HBcAg produced from pFS14 has been demonstrated by characteristic sedimentation properties in sucrose gradients, by banding in cesium chloride density gradients, and by electron micrography (data not shown). Immunization with recombinant HBV core particles purified by cesium chloride density gradient centrifugation induced a protective immune response against challenge with woodchuck hepatitis virus (WHV) in two of three woodchucks (F. Schödel et al., in prep.). We therefore assume that amino acid sequences conserved between HBV and WHV mediate protective immunity (for an amino acid comparison between hepatitis virus core proteins, see Schödel et al. 1989). An interesting candidate is the region between amino acids 120 and 140 of HBcAg, which is highly conserved between WHV and HBV, encompasses two T-cell sites recognized in the murine model by mice of H-2^b and H-2^s haplotype, and is also immunogenic at the B-cell level as a free peptide (Milich et al. 1987a).

Construction of Core/Pre-S2 fusions

Amino-terminal Fusions

We first set out to construct expression vectors that would direct the synthesis of core proteins with the pre-S2 *ayw* (amino acids 133-140) sequence fused to the amino termi-

nus of HBcAg. fused the sequ dem repeat to in a *lac* promo could not be c and pre-S2/coi

Internal Fusion

A recent mod RNA virus nur that the site c hepadnavirus, tional fusion c quence, we i synthetic oligc GTA CTT CC coding for Hf unique XhoI s and colonies body (Okamc from positive the resultant and lysed Jlv proteins on a behavior of p shown). Thus tion. Whether is unresolved 4408 or 552t (120-145) (p HBcAg do not ticles is inter

Carboxy-ter

As a third sit HBcAg. For 461 (for nuc ment clonec al. 1989). T placed by a 155 is there placed by t (Fig. 1). Usi inserted tha 156. In the tre me carb spective us inserted a GTT CGT

nus of HBcAg. Using 5'-degenerate oligonucleotides in polymerase chain reactions, we fused the sequence coding for amino acids 133-140 of the HBV pre-S2 region as a tandem repeat to the 5' end of the core gene in a *trc* promoter vector and as a single repeat in a *tac* promoter vector. With both vectors, stable core or pre-S2 reactive fusion protein could not be detected in the appropriate *E. coli* hosts (not shown), although promoter and pre-S2/core gene sequences were not mutated, as verified by dideoxy sequencing.

Internal Fusion

A recent model for the HBcAg particle structure, based primarily on homologies with RNA virus nucleocapsid particles for which structural information is available, suggests that the site of a 39-amino-acid insertion in the core protein of duck HBV, a related hepadnavirus, is exposed on the surface (Argos and Fuller 1988). To achieve translational fusion of the pre-S2 *ayw* (amino acids 133-143) sequence within the HBcAg sequence, we digested pFS14 partially with *Xba*I, inserted a pair of complementary synthetic oligonucleotides (coding strand 5'-CTA GGG ACC GCG TGT TCG TGG TCT GTA CTT CCC GGC TCG GAG-3') coding for the pre-S2 sequence 3' of the sequence coding for HBcAg up to amino acid position 82, and introduced at the same time a unique *Xho*I site at the 3' end of the insert (pFS14PS1, Fig. 1). JM105 was transformed, and colonies were screened for expression of pre-S2 epitopes using a monoclonal antibody (Okamoto et al. 1985) kindly provided by M. Mayumi. Plasmid DNA was isolated from positive colonies, the inserted sequence was verified by dideoxy sequencing, and the resultant fusion protein was analyzed by immunoblotting (Fig. 2, left). We induced and lysed JM105 (pFS14PS1) under nondenaturing conditions and separated soluble proteins on a Sepharose 4B column. Anti-core reactivity identical to the chromatographic behavior of particles expressed from pFS14 was found only in the exclusion volume (not shown). Thus, we assume that the internal fusion does not interfere with particle formation. Whether hybrid particles from pFS14PS1 carry the pre-S2 epitopes on their surface is unresolved. Raw bacterial lysates do not react with anti-pre-S2 monoclonal antibodies 4408 or 5520 (Okamoto et al. 1985), but these lysates do react with rabbit anti-pre-S2 (120-145) (provided by R. Neurath). As expected, most monoclonal antibodies against HBcAg do not recognize products from pFS14PS1, since the major epitope on core particles is interrupted by the insertion.

Carboxy-terminal Fusion

As a third site for fusing pre-S2 sequences to HBcAg, we chose the carboxyl terminus of HBcAg. For this purpose, a unique *Ap*I site was introduced at HBV nucleotide position 461 (for nucleotide position references, see Schödel and Will 1989) in an HBV subfragment cloned into a plasmid vector suitable for site-directed mutagenesis (Stanssens et al. 1989). The T residue at position 463 of the authentic *ayw* HBV sequence was replaced by a G, resulting in plasmid pMHM7. The serine at HBcAg amino acid position 155 is thereby changed to an alanine. The core gene *Bgl*II fragment of pFS14 was replaced by the corresponding *Bgl*II fragment from pMHM7 to give plasmid pFS14apa (Fig. 1). Using the unique *Ap*I site in pFS14apa, heterologous gene fragments can be inserted that result in fusion proteins with HBcAg carboxy-terminal to amino acid position 156. In these fusion proteins, the arginine-rich nucleic-acid-binding domain at the extreme carboxyl terminus of HBcAg is deleted, which may be an advantage for prospective use as a vaccine, and seems not to interfere with particle formation. We have inserted a pair of synthetic oligonucleotides (coding strand 5'-CCG GAC CCG CGT GTT CGT GGT CTG TAC TTC CCG GCT TA-3') specifying again an HBV pre-S2 se-

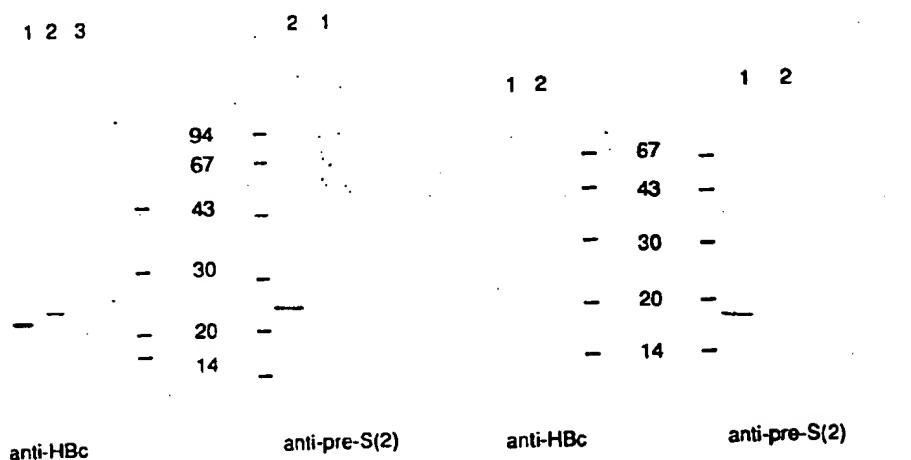


Figure 2
Immunoreactivity of hybrid HBc/pre-S2 fusion proteins from pFS14PS1 (left) and pFS14PS2 (right) expressed in *E. coli* JM105. Bacterial overnight cultures without plasmid (JM105) or with the indicated plasmid were diluted 1:10 in Standard 1 medium, grown to a density of approximately 0.8 OD₆₀₀ and induced at 37°C by 1 mM IPTG for 1 hr. For conditions of electrophoresis (on 15% SDS-PAGE) and immunoblotting, see Schödel and Will (1989). Anti-HBc is a polyclonal rabbit serum raised against a purified MS2-HBcAg fusion expressed in *E. coli*, and anti-pre-S2 is a monoclonal antibody (hybridoma 5520) kindly provided by M. Mayumi (Okamoto et al. 1985). The left blot in the left panel is labeled with ¹²⁵I-protein A. All of the other blots are with alkaline phosphatase-coupled goat anti-mouse or anti-rabbit IgG (H+L) (Medac) using NBT/BCIP as substrate.

quence (133–143), resulting in expression plasmid pFS14PS2 (Fig. 1). A fusion protein with core and pre-S2 reactivity of the expected molecular weight was observed in immunoblots from induced JM105 (pFS14PS2) raw extracts (Fig. 2, right). Nondenatured proteins of the JM105 (pFS14PS2) whole-cell bacterial lysate separated on Sepharose 4B again displayed anticore reactivity only in the void volume identical to that of pFS14 and pFS14PS1 (not shown), which we interpret as evidence for particle formation. The fused pre-S2 epitopes are also detected by anti-pre-S2 monoclonal antibodies 4408 and 5526, as well as rabbit antipeptide pre-S2 (120–145) in nondenaturing ELISAs of bacterial lysates, strongly suggesting their accessibility on the particle surface.

SUMMARY AND CONCLUSION

We have constructed a prokaryotic expression vector directing the synthesis of HBcAg of the authentic *ayw* sequence (pFS14) in *E. coli*. Gene fragments coding for immunodominant conserved B-cell epitopes of the pre-S2 region of HBV surface antigens were fused to the amino terminus, to internal HBcAg sequences, and to the carboxy terminus of HBcAg. Amino-terminal fusions did not lead to expression of stable fusion proteins. Both internal and carboxy-terminal fusions, the latter also removing the HBcAg nucleic-acid-binding region, led to expression of stable fusion proteins with dual reactivity in immunoblots. B-cell epitopes fused to the carboxyl terminus of HBcAg were accessible to antibodies in nondenaturing assays and are thus probably located at the surface of HBcAg particles. The immunoreactivity of purified hybrid HBcAg/pre-S2 particles is cur-

rently being tested for cell epitopes and accessibility to be determined. The epitopes at the surface and of study in

ACKNOWLEDGEMENTS

We thank R. laboratory was Sander Stittur

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rently being tested, and it will be especially interesting to determine whether the fused B-cell epitopes will acquire T-cell-independent immunogenicity. In addition, size limits and accessibility of different internal inserts as well as carboxy-terminal fusions will be determined. The existence of HBcAg hybrids with the same or different heterologous epitopes at separate amino acid positions raise the possibility of creating mixed particles and of studying their immunogenicity.

ACKNOWLEDGMENTS

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Expression of HIV *gag* and *env* B-cell Epitopes on the Surface of HBV Core Particles and Analysis of the Immune Responses Generated to Those Epitopes

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An investigation into the presence of T-cell epitopes within the hepatitis B virus (HBV) has led to the identification of a potential universal carrier molecule in the form of the hepatitis B core antigen (HBcAg) particle. The HBc particle, which exhibits unique immunological characteristics, can be engineered to carry multiple, heterologous epitopes from other pathogens that may ultimately serve as vaccine candidates. We have selected human immunodeficiency virus (HIV) *gag* and *env* sequences as candidate B-cell epitopes to be positioned on the surface of HBc particles and expressed as fused proteins. The DNA constructs were prepared in a manner that would not alter the inherent T-cell functions and particulate nature of the HBc particles.

Unique Immunological Characteristics of HBc Particles

The use of hepatitis B particulate antigen as a T-cell carrier protein is not unique to HBc. The hepatitis B surface antigen (HBsAg) has been utilized as a T-cell carrier protein for heterologous B-cell epitopes. A comparison of the humoral response generated to HBcAg versus that observed with HBsAg suggests that the core antigen of HBV, rather than the surface antigen, represents a more potent immunogen and hence a better T-cell carrier. The development of antibody to the HBsAg is not achieved by all individuals infected with HBV; however, virtually all of those infected with the virus develop antibody to HBc. A comparison of HBc with HBs in seven congenic murine strains demonstrated that HBc could elicit high antibody titers in all strains tested, whereas HBs was reactive in only five of the seven, with the generation of low-to-moderate antibody titers (Millich and McLachlan 1986).

The ability of HBcAg to function as a T-cell-independent antigen has also been dem-

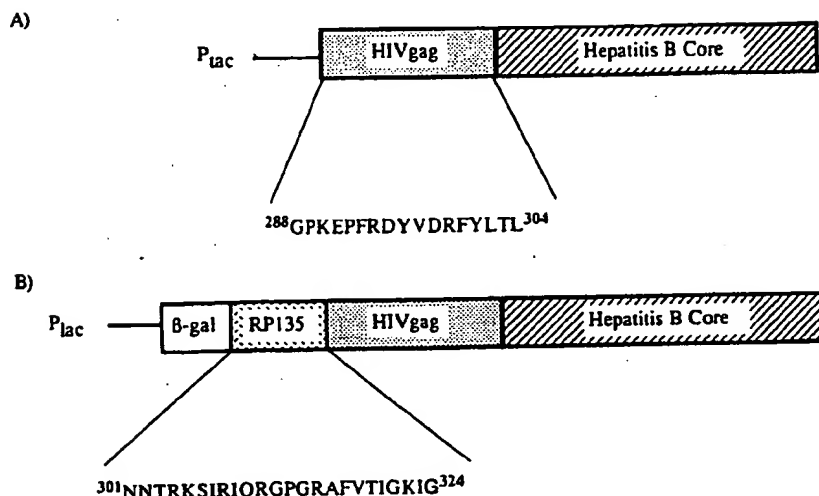


Figure 1

(A) An HIV *gag* sequence (288–304), inserted upstream of the HBc-coding sequence cloned in vector pKK223-3, was expressed as a fusion protein in *E. coli*. The synthetic DNA oligomer, complementary to the selected amino acid sequence, was inserted at *EcoRI* site in the multicloning site of vector. (B) An HIV *env* sequence (RP135) was inserted upstream of the HIV *gag*/HBc done in plasmid pBS at the *PstI*-*Bam*HI insertion site of the vector.

onstrated with the particulate form of HBcAg. Immunization of a euthymic strain of mouse and its athymic counterpart with particulate HBc resulted in antibody production in both strains, demonstrating that HBcAg could function as a T-cell-independent antigen (athymic), as well as a T-cell-dependent antigen (euthymic) (Milich and McLachlan 1986). This unique immunological characteristic was not due to the particulate nature of HBc alone in that HBs particles were unable to elicit antibody by a T-cell-independent pathway. Additionally, it had been demonstrated that attachment of a hapten (DNP) to HBc through chemical coupling resulted in the same T-cell independence exhibited by HBc (Milich et al. 1988). All of these immunological features exhibited by particulate HBc demonstrate its uniqueness as a T-cell carrier molecule.

DISCUSSION

Expression of a T/B-cell Epitope from the HIV *gag* Region on the Surface of HBc Particles

To determine if HBc particles can act as a carrier for a foreign B-cell epitope, we selected the HIV *gag* region of HIV-1 for investigation. Our intent was to select peptide sequences that provide T- and B-cell functions and present them on the surface of HBc particles to take advantage of the unique cellular mechanisms that have been defined for HBc. A peptide sequence (288–304), predicted by standard algorithms to contain both a T- and B-cell epitope within a limited amino acid region (17-mer) (Coates et al. 1987), was inserted upstream of the HBc coding sequence (AM6) (Moriarty et al. 1981) in the *Escherichia coli* expression vector pKK223-3 (Pharmacia). A DNA oligomer representing the nucleotide complement of the 288–304 amino acid sequence (Fig. 1A) was inserted at the *EcoRI* site of the vector. This particular peptide was weakly immunogenic

Table 1
Determining the A
of the HIV Epitope

Antibody used to
capture particles:

Labeled antibody
to probe for antigen

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Purified HIVgag/t

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Table 1
Determining the Antigenicity of the Purified pKK/HIVgag/HBc Particles and Confirming the Location of the HIV Epitope on the Surface of HBc

Antibody used to capture particles:	-	-	-	-	Anti-288-304
Labeled antibody used to probe for antigen:	Anti-HBc	Anti-288-304	Anti-p24	Anti-HIV lysate	Anti-HBc
Purified HBc	1.618	.019	n.d.	.011	.012
Purified HIVgag/HBc	1.626	1.920	1.852	.771	1.827

Numbers represent OD values recorded at 490 nm. The same concentration of antigen was used for both preparations (100 ng). The antibodies used to probe for antigen were either rabbit polyclonal antibodies generated to peptides and whole proteins or mouse monoclonal antibodies specific for HBc. n.d. indicates not determined.

on its own; it was able to raise antibody in both rabbits (after three immunizations) and mice (BALB/c), demonstrating that a T-cell site was contained within the 288-304 sequence that was capable of eliciting antibody in the animals tested. In addition, it should be noted that a human serum sample positive for HIV-1 was capable of reacting with this peptide in ELISA. Also, antibodies raised in rabbits to both a recombinant p24 preparation (Centocor) and an HIV lysate were able to react with the 288-304 peptide, suggesting that it may contain a native B-cell site.

A DNA construct (pKKHIVgag/HBc) was used to express the hybrid protein in *E. coli*, and the fused product was detected by Western blot analysis with antibody specific for denatured HBc and an anti-p24 rabbit antibody. A protein with the predicted size of 23,000 M_r (the HIV 17-mer plus linker sequences, six precore amino acids, and the full HBc sequence [21,000 M_r]) was demonstrated. The material was purified to homogeneity by means of gel filtration, ion-exchange chromatography, and hydroxyapatite binding. The material obtained after the purification procedure was analyzed by ELISA and determined to be particulate (reactive with anti-HBc), and the HIV gag epitope was positioned on the surface of the particles (anti-288-304 could capture the particles) (Table 1). Purified HIV gag/HBc particles were reactive with anti-HBc, anti-288-304, anti-p24, and anti-HIV lysate. The particulate nature of the material was subsequently confirmed by electron microscopy, where the hybrid particles were found to be identical in shape, although slightly larger in size, than native HBc particles (31 nm vs. 27 nm).

Immune Response in Mice Generated to the Purified HIVgag/HBc Particles

The purified HIV gag/HBc particles were used to immunize mice, previously identified as nonresponders to peptide 288-304 (B10 and B10.S). The peptide was presented both linked to keyhole limpet hemocyanin (KLH) and on the surface of HBc particles (HIV gag/HBc particles) to compare the immune response generated to the attached epitope as a function of the carrier molecule (Table 2). The antibody responses generated were evaluated by reactivity with peptide 288-304 (antipeptide) and a recombinant p24 protein (anti-gag). A primary immunization was administered at day 0, followed by a boost at day 30. The animals were bled at days 24 and 44. Peptide 288-304, administered alone, is nonreactive in both strains of mice even after a boost. The delivery of peptide 288-304 linked to KLH resulted in antipeptide titers at day 24 in both strains; however, there was no anti-gag response at the same time point. The presentation of peptide on the surface of HBc particles resulted in a response similar to the antipeptide response

Table 2
Antibody Titers Generated in B10 Mice Strains to HIVgag Epitopes

Immunogen	Anti-peptide		Anti-gag	
	B10	B10.S	B10	B10.S
288-304				
primary	0	0	0	0
boost	0	0	0	0
288-304-KLH				
primary	1,280	320	0	0
boost	80,000	1,280	20,000	320
288-304/HBc				
primary	1,280	1,280	1,280	1,280
boost	>80,000	5,000	>80,000	5,000

The concentrations of immunogen used were 100 µg peptide, 20 µg of peptide linked to KLH, and 10 µg of HIVgag/HBc particles. The values represent reciprocal titers. Animals were bled 24 days after a single immunization (primary) and 14 days after a boost at day 30 (boost).

recorded for the peptide-KLH preparation at day 24; however, an anti-gag response was generated after only a single immunization, which increased significantly after the boost. The presentation of the HIV gag epitope on the surface of HBc particles was more immunogenic than the peptide linked to KLH (higher antibody titers) and elicited a stronger and more rapid anti-native response. Further immunological analysis in higher animals should confirm the effectiveness of HBc as a carrier for foreign epitopes.

Insertion of an HIV *env* Sequence Upstream of the HBc and HIVgag/HBc Sequences

The T-cell help provided by HBc for the carried B-cell epitope is nonspecific with regard to the foreign sequence. In an attempt to provide specific T-cell help to a carried B-cell epitope, an HIV envelope sequence identified as a neutralizing epitope (RP135) (Rusche et al. 1988) was inserted upstream of the HIV gag/HBc fusion protein. We selected a gag T-cell sequence over those predicted for the envelope region in an effort to determine if the HIV gag region influences the immune response generated to the HIV envelope in a manner similar to that demonstrated with HBV (Milich et al. 1987). The internal viral protein of HBV (HBc) regulates the immune response generated to the HBV envelope proteins (HBs). Insertion of a B-cell epitope from the HIV envelope upstream of an HIV gag T-cell site, and positioning both on the surface of HBc particles, has been accomplished. A DNA oligomer complementary to the RP135 sequence was inserted at the *Pst*I-*Bam*HI site of plasmid pBS (Stratagene) in which the HIV gag/HBc-coding sequence was downstream (Fig. 1B). The final product is a fusion protein consisting of five β -galactosidase amino acids, linker sequences, the HIV gag 17-mer, six precore residues, and the entire HBc sequence. The predicted size for the fusion protein is 26,000 M_r. Western blot analysis revealed a protein species migrating at the predicted molecular weight for the fusion protein recognized by antibody specific for HBc and the RP135 determinant. ELISA confirmed the presence of particles (reactive with anti-HBc) and the surface location of the RP135 envelope sequence (anti-RP135 can capture the hybrid particles). These particles are currently being purified for immunological analysis.

SUMMARY

The ability of HE particle formation manner versus anti-gag antibody (B10 and B10.S) dependence when presence of HIV gag will be determined by gal/RP135/HIV.

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SUMMARY

The ability of HBc to carry epitopes (HIV *gag* and *env*) on its surface, while maintaining particle formation, has been demonstrated. The presentation of B-cell epitopes in this manner versus that observed when coupled to KLH generated higher antipeptide and anti-*gag* antibodies in two strains of mice that were nonresponders to the peptide alone (B10 and B10.S). The ability of the HIV *gag* epitopes to demonstrate T-cell independence when presented on the surface of HBc particles remains to be elucidated. The ability of HIV *gag* to influence the immune response generated to the envelope of HIV will be determined with the hybrid particles expressed with the DNA construct β -gal/RP135/HIV *gag*/HBc.

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